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IV

(54) Title: A METHOD FOR OBTAINING STRUCTURAL INFORMATION CONCERNING AN ENCODED MOLECULE AND METHOD FOR SELECTING COMPOUNDS

(57) Abstract: In one aspect, the present invention relates to a method for obtaining structural information about an encoded molecole call. The encoded molecole may be produced by a reaction of a plurality of chemical entities and may be capable of being connected to us identifier oligonuclocule containing codons informative of the identity of the chemical entities which have participated in the formation of the encoded molecule. In a acretin term bodiment, primers are designed complementary to the codons appearing on the IS identifier oligonuclocitide, and the presence absence or relative abundance of a codon is evaluated by mixing a primer with the idendifier oligonuclocitide, and the presence of a polymentees and substrate (deaxy) ribonuclocitide triphosphates measuring the extension of entire oligonuclocities in the presence of a polymentee and substrate (deaxy) ribonuclocities triphosphates measuring the extension creation. In another aspect, the invention provides a method for selecting compounds which binds to a target. More specifically, the invention relates to a method in which a larget associated with an oligonuclocitie initially is mixed with a library of complexes, each invention relates to a method in which a target associated with an oligonucleotide initially is mixed with a library of complexes, each the target oligonucleotide is coupled to the identifier oligonucleotide of complexes having a display molecule with affinity towards complex comprising a display molecule and an oligonucteotide identifying said display molecule. Next, due an increased proximity, the target. In a final stage the coupled nucleotides are analysed to deduce at least the identity of the display molecule.

WO 2005/026387

PCT/DK2004/000630

A method for obtaining structural information concerning an encoded molecule and method for selecting compounds

Technical Field of the Invention

- tural information about an encoded molecule. The encoded molecule may be of being connected to an identifier oligonucleotide containing codons informaproduced by a reaction of a plurality of chemical entitles and may be capable ive of the identity of the chemical entities which have participated in the for-In one aspect, the present invention relates to a method for obtaining struc-
- ated by mixing a primer with the identifier oligonucleotide in the presence of a signed complementary to the codons appearing on the identifier oligonucleopolymerase and substrate (deoxy)ribonucleotide triphosphates and measurtide, and the presence, absence or relative abundance of a codon ls evalu∗ mation of the encoded molecule. In a certain embodiment, primers are de-9
 - comprising a display molecule and an oligonucleotide identifying sald display method for selecting compounds which binds to a target. More specifically, the invention relates to a method in which a target associated with an ollgonucleotide initially is mixed with a library of complexes, each complex ing the extension reaction. In another aspect, the invention provides a ñ
- coupled to the Identifier oligonucleotide of complexes having a display molecule with affinity towards the target. In a final stage the coupled nucleotides nolecule. Next, due an increased proximity, the target ollgonucleotide is are analysed to deduce at least the identity of the display molecule. 2
- A method for obtaining structural information concerning an encoded 22

The below paragraphs up to the section entitled "Method for identifying a display molecule" relate to the first aspect of the invention

structural information about an encoded molecule. The encoded molecule is usually produced by a process that comprises the reaction of a plurality of The first aspect of the present invention relates to a method for obtaining ဓ

PCT/DK2004/000630

molecule. The structural information obtained by the present method may be programmed in an identifier oligonucleotide which is attached to the encoded used to obtain the entire structure of the encoded molecule or a part thereof. chemical entities. The synthesis of the encoded molecule is recorded or

Background of the Invention

The generation of molecules carrying new properties remains a challenging allow a more efficient generation and screening of a huge number of moletask. Recently, a number of procedures have been suggested that should

molecules other than natural biopolymers and a coupling of the molecules to respective templates or identifier parts containing information about the reaccules. The approach taken may involve the encoding and/or templating of proaches allow the researcher to generate and screen a huge number of tants that have participated in the formation of the molecule. These ap-은 5

molecules at the same time.

in which one part of the bifunctional molecule comprises an encoded part and the other part of the molecule contains an identifying part. The identifying part In US 5,723,598 it is suggested to prepare libraries of bifunctional molecules,

mix method, which involves the initial reaction between a nascent bifunctional molecule and a range of different reactants in separate compartments at one end of the nascent bifunctional molecule and a corresponding range of identifier unit oligonucleotides (codons) and the other end. Subsequently, the conreactants that have been involved in the synthesis of the encoded molecule. The libraries of bifunctional molecules are generally prepared by a split-andis segregated into codons, i.e. a stretches of nucleotides, which codes for 22 8

molecules, a partitioning with respect to affinity towards a target is conducted and the identifier part of the bifunctional molecule is decoded to establish the chemical structure of the compounds in the library that is likely to be a ligand sponding codons. Following the generation of a library of the bifunctional ဗ္က

tents of the compartments are mixed and the mixture is disposed in separate

compartments and reacted again with another range of reactants and corre-

PCT/DK2004/000630 WO 2005/026387

initially are amplified by PCR. The PCR product is subsequently incorporated to the target. The decoding step implies that the identifier oligonucleotides in to a suitable vector which is transformed to a host organism, usually E. coli. Following the incubation of the E. coli, colonies are picked and se-

quenced. 2

approach stipulated above. The approach is based on the same split-and-mlx end a chemical reactive site and dispersed throughout the stand a plurality of codon regions, each of said codon regions in turn specifying different codons. Halpin and Harbury have in WO 00/23458 suggested an improvement to the all the strands are pooled and subjected to a second partitioning based on a at the chemical reaction sites with specific selected reagents. Subsequently, second codon region. The split-and-combine method is conducted an appro-Separately, each of the strands, identified by a first codon region, is reacted . thetic steps. A plurality nucleic acid templates are used, each having at one strategy for synthesis of combinatorial libraries comprising two or more synpriate number of times to produce a library of typically between $10^3\,\mathrm{and}~10^6$ different compounds. The decoding is performed utilizing the process depicted above. 9 5

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short, a template segregated into a plurality of codons and a plurality of buildtemplate oligonucleotides initially are amplified by PCR. The PCR product is Recently, a new method for encoding molecules has been suggested, which blocks are annealed together and the chemical entities are subsequently reselections have been performed, the template must be decoded to establish close methods for preparing virtually any molecule connected to a template acted to form the molecule. However, after a sufficient number of rounds of initially provided. Under hybridisation conditions, the template and building can be performed in a single "pot". WO 02/00419 and WO 02/103008 dising blocks comprising a transferable chemical entity and an anticodon are coding for chemical entities which have reacted to form the molecule. In the identity of the encoded molecule. The decoding step implies that the 22 ജ

WO 2005/026387

subsequently incorporated in to a suitable vector which is transformed to a host organism, usually E. coli. Following the incubation of the E. coli, colonies are picked and sequenced.

- In an aspect of the invention, it is the object to facilitate the decoding of the coding oligonucleotide in order to obtain at least partial structural information of the encoded molecule being a ligand towards a target. In another aspect of the invention, it is desired to obtain information about which chemical entities that result in encoded molecules successful in a selection process. Such chanical entities may be used in the formation of open desired in the formation.
 - chemical entities may be used in the formation of a second generation library.

Summary of the Invention

The first aspect of the present invention concems a method for obtaining structural information about an encoded molecule produced by a process comprising reaction of a plurality of chemical entities, said encoded molecule being capable of forming part of a complex also comprising an identifier oligonucleotide containing codons informative of the identity of chemical entities which have participated in the formation of the encoded molecule, the

cleotide, subjecting the mixture to a condition allowing for an extension reaction to occur when the primer is sufficient complementary to a part of the identifier oligonucleotide, and evaluating, based on measurement of the extension reaction, the presence, absence, or relative abundance of one or 25 more codons.

The method according to the invention may be performed on a single identifier oligonucleotide or a composition of identifier oligonucleotides to obtain structural information about the encoded molecule or a composition of encoded molecules, respectively, that have been attached to the identifier oli-

gonucleotide(s)

PCT/DK2004/000630

WO 2005/026387

2

PCT/DK2004/000630

A single identifier may be analysed using the above method to verify the incorporation into the encoded molecule of one or more chemical entities or to deconvolute the identity of the entire encoded molecule. A composition of two or more identifier oligonucleotides generally results from a selection

process, i.e. a process involving subjecting a library of different complexes to a condition partitioning the composition from the remainder of the library. Usually the partitioning condition includes an affinity assay in which the library of complexes is contacted with a target and the Identifier oligonucleotides of the binding complexes are harvested.

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The conditions allowing for an extension reaction to occur may be selected from a enzymatic or chemical means. Suitably, the condition involves one or more enzymes. In a certain embodiment of the invention, the condition which allows for an extension reaction to occur includes a polymerase or a ligase

as well as suitable substrates for the enzyme used. Preferred is a polymerase together with a blend of (deoxy)ribonucleotide triphosphates. Suitably, the blend include one or more of dATP, dGTP, dCTP, and dTTP.

A library of complexes can have any appropriate size. Typically, the size is

- above 10³, typically above 10⁶ different complexes. An effective, extensive, and rapid decoding is therefore desirable. The method of the present invention may be used at various stages of the process of finding a ligand to a certain target. As examples, the method of the invention may be used for controlling the quality of a starting library. The information acquired may be used
- 25 to verify which codons being present, absent, and, in some embodiments, also the relative abundance. Thus, the method of the Invention delivers a reliable picture of the process which has produced the library. If, for some reason, a chemical entity has not been incorporated into the encoded molecules, the absence of a codon for this chemical entity will in certain embodiments of
- 30 the invention indicate this fact.

chemical entities that have been used in the synthesis of encoded molecules having an affinity towards the target. In the event the selection has been suf-Another example of the use of the present method is following the selection. After the selection has been performed the codon profile is indicative of the

- ture-activity-relationship (SAR). If the selection process has not narrowed the molecules after the selection, which gives important information to the strucstructure of binding encoded molecules. Alternatively, it may be possible to ficiently effective it may be possible directly to deduce a part or the entire deduce a structural unit appearing more frequently among the encoded œ.
 - tion library chemical entities which have not been involved in the synthesis of then be subjected to more stringent selection conditions to allow only the eneration library may also be spiked with certain chemical entities suspected of encoded molecules that have been successful in the selection may be omitcoded molecules with a higher affinity to bind to the target. The second generation library may be contemplated. In the formation of the second generated, thus limiting the size of the new library and at the same time increasing the concentration of binding complexes. The second generation library may size of the library to a manageable number, the formation of a second gen-9 5
- certain successful chemical entities may be obtained from the SAR. The use increasing the performance of the final encoded molecule. The indication of in a second generation library of chemical entities, which have proved to be interesting for further investigation in a preceding library, may thus entail a shuffling with new chemical entities that may focus the second generation library in a certain desired direction. 8

contain the same codon. Thus, following the formation and selection of a first, The relative abundance of codons may make it possible to decode a plurality of identifiers simultaneously, even in the case when two or more identifiers second or further generation library, the identity of binding encoded mole-

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cules may be partly or entirely deconvoluted by the present method ဓ္က

PCT/DK2004/000630 WO 2005/026387

vention n is an integer independently selected from of 2 to 8. It may be pregonucleotides having n codon positions each. In a certain aspect of the in-In a practical approach the library comprises complexes with identifier oli-

- erred that n is constant among all the complexes in the library to facilitate the codon position or may be constant among the various codon positions. It may decoding process. Each of the codons in a certain position is in an aspect of the invention selected from a set of m different codons. m may vary for each be preferred in some embodiments to have all the codons in each position selected from the same set of m codons. However, in other embodiments, Ġ
- especially such involving hybridisation in the recognition between the codon and the anticodon, it may be preferred that all the codons are different. ₽.

of the set. In some embodiments of the invention, a set of primers comprising Preferably, any member of the codon set differs from any other codons in the ber of the codon set differs with at least two nucleotides nucleotide positions set with the identity of at least one nucleotide, i.e. at least one nucleotide pogeneral, it is desired to maximize the differences between individual codons sition occurs. In some aspects of the invention it is preferred that any memfrom any other member of the set to increase the fidelity of the method. In a sequence of complementing the set of codons are prepared 5 8

In a preferred aspect of the method a framing sequence is related to each of tions the reaction of a chemical entity in the synthesis history of the encoded the n codon positions in a particular complex, said framing sequence posi-

- ent codons and the set of n different framing sequences is prepared. The n x molecule. Typically, the framing sequence is identical among the complexes primers fully or in part complementing any combination of the set of m differfor each of the reaction rounds and is selected from a group of n different nucleotide sequences. In a certain aspect of the invention n x m different 23
- m primers may be used in separate compartments to reveal the identity of a chemical entity as well as the point in time of the synthesis of the encoded molecule is has reacted. ജ

chemical entities utilized in the formation of an encoded molecule or a composition of encoded molecules, wherein in separate compartments, n x m in a particular aspect, the invention concerns a method for identifying the

- composition from the remainder of the library, subjected to a mixture of polymerase and substrate (deoxy)ribonucleotide triphosphates under conditions primers individually are mixed with an aliquot of a composition obtained by allowing for an extension reaction to occur when a primer is sufficient comsubjecting a library of different complexes to a condition partitioning said Ŋ
 - aliquot, and evaluation, based on measurement of the extension reaction, the plementary to a part of one or more identifier oligonucleotides present in the presence, absence, or relative abundance of one or more codons in each 9
- primers, a polymerase, a composition of (deoxy)ribonucleotide triphosphates sufficiently complementary to codons appearing on the identifier oligo nucleoformative of the identity of the chemical entities which has participated in the The invention also concerns a set comprising a collection of oligonucleotide and an identifier oligonucleotide, said oligonucleotide comprising codons in-(dNTPs), and a library of complexes composed of a display molecule part formation of the display molecule, wherein the oligonucleotide primers are lides in the library to allow for an extension to occur. 5 2

Detailed Description of the Invention

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uniquely, i.e. in a library of complexes a particular identifier is capable of dis-The complex comprises an encoded molecule and an identifier oligonucleotide. The identifier comprises codons that identify the encoded molecule. Preferably, the identifier oligonucleotide identifies the encoded molecule

tinguishing the molecule it is attached to from the rest of the molecules.

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WO 2005/026387

PCT/DK2004/000630

other or through a bridging molety. In one aspect of the invention, the bridg-The encoded molecule and the identifier may be attached directly to each ing moiety is a selectively cleavable linkage.

- The sequence of each codon can be decoded utilizing the present method to dentifier comprises more than one codon, each member of a pool of chemiferred aspect the identifier oligonucleotide comprises three or more codons. identify reactants used in the formation of the encoded molecule. When the The identifier oligonucleotide may comprise two or more codons. In a preß
- cal entities can be identified and the order of codons is informative of the synthesis step each member has been Incorporated in. 9

In a certain embodiment, the same codon is used to code for several different chemical entities. In a subsequent Identification step, the structure of the en-

- protection groups, etc. In another embodiment, the same codon is used for a nature, a certain attachment chemistry etc. In a preferred embodiment, howcoded molecule can be deduced taking advantage of the knowledge of diferent attachment chemistries, steric hindrance, deprotection of orthogonal group of chemical entities having a common property, such as a lipophilic र्ठ
 - appear on the identifier oligonucleotide coding for another chemical entity. In a practical approach, for a specific chemical entity, only a single combination geous to use several codons for the same chemical entity, much in the same ever, the codon is unique i.e. a similar combination of nucleotides does not of nucleotides is used. In some aspects of the invention, it may be advanta-2
 - way as Nature uses up to six different codons for a single amino acid. The wo or more codons identifying the same chemical entity may carry further nformation related to different reaction conditions. 22

The sequence of the nucleotides in each codon may have any sultable

length. The codon may be a single nucleotide or a plurality of nucleotides. In some aspects of the invention, it is preferred that each codon independently ജ

comprises four or more nucleotides, more preferred 4 to 30 nucleotides. In some aspects of the invention the lengths of the codons vary,

ess and to increase the ability of the primer to discriminate between codons it certain number of nucleotides in the codon, it is generally desired to optimize A certain codon may be distinguished from any other codon in the library by only a single nucleotide. However, to facilitate a subsequent decoding procdeotide combinations exist in which two or more mismatches appear. For a the number of mismatches between a particular codon relative to any other is in general desired to have two or more mismatches between a particular example, if a codon length of 5 nucleotides is selected, more than 100 nucodon and any other codon appearing on identifier oligonucleotide. As an codon appearing in the library.

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ing sequence may have any suitable number of nucleotides, e.g. 1 to 20. Alformed, the identifier may comprise further codons, such as 3, 4, 5, or more are separated from a neighbouring codon by a framing sequence. The framcodons. Each of the further codons may be separated by a suitable framing sequence. Preferably, all or at least a majority of the codons of the identifier ranged in sequence, i.e. next to each other. Two neighbouring codons may be separated by a framing sequence. Depending on the encoded molecule ternatively, codons on the identifier may be designed with overlapping se-The identifier oligonucleotide will in general have at least two codons arquences. 5 8

The framing sequence, if present, may serve various purposes. In one setup conditions in the synthesis history of the encodedmolecule. The framing sequence may also or in addition provide for a region of high affinity. The high comprises information which positions the chemical entity and the reaction affinity region may ensure that a hybridisation event with an anti-codon will of the invention, the framing sequence identifies the position of the codon. Usually, the framing sequence either upstream or downstream of a codon ജ

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PCT/DK2004/000630 WO 2005/026387

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occur in frame. Moreover, the framing sequence may adjust the annealing temperature to a desired level. A framing sequence with high affinity can be provided by incorporation of one affinity, such as 2'-O-methyl substitution of the ribose moiety, peptide nucleic sine. Alternatively, or in addition, the framing sequence may be subjected to backbone modification. Several back bone modifications provides for higher base. Examples of nucleobases having this property are guanine and cytoor more nucleobases forming three hydrogen bonds to a cognate nucleo-2

acids (PNA), and 2'-4' O-methylene cyclisation of the ribose molety, also referred to as LNA (Locked Nucleic Acid). 9

The sequence comprising a codon and an adjacent framing sequence has in a certain aspect of the invention a total length of 11 nucleotides or more, preferably 15 nucleotides or more. A primer may be designed to complemenwell as the position said chemical entity has in the entire synthesis history of of an extension reaction under conditions allowing for such reaction to occur lary to the codon sequence as well as the framing sequence. The presence is indicative of the presence of the chemical entity encoded in the codon as 5

the encoded molecule. გ. The identifier may comprise flanking regions around the coding section. The flanking regions can also serve as priming sites for amplification reactions, such as PCR or as binding region for oligonucleotide probe. The Identifier

may in certain embodiments comprise an affinity region having the property of being able to hybridise to a building block. 22

It is to be understood that when the term Identifier oligonucleotide is used in the present description and claims, the identifier oligonucleotide may be in the sense or the anti-sense format, i.e. the identifier can be a sequence of codons which actually codes for the encoded molecule or can be a sequence

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complementary thereto. Moreover, the identifier may be single-stranded or double-stranded, as appropriate.

pected of having an effect on a target. When the target is of pharmaceutical candidates with a tag, e.g. a nucleic acid tag identifying each possible drug The complex may be formed by tagging a library of different possible drug importance, the encoded molecule is generally a possible drug candidate. The encoded molecule part of the complex is generally of a structure ex-Ŋ

molecule. Optionally, this reaction product may be post-modified to obtain the candidate. In another embodiment of the invention, the molecule formed by a the cleavage of one or more chemical bonds attaching the encoded molecule final molecule displayed on the complex. The post-modification may involve variety of reactants which have reacted with each other and/or a scaffold to the identifier in order more efficiently to display the encoded molecule. è

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connection to another reactive group positioned on a chemical entity, thereby generating an addition to the original scaffold. A second chemical entity may The formation of an encoded molecule generally starts by a scaffold, i.e. a entities may be involved in the formation of the final reaction product. The reactive group incorporated by the first chemical entity. Further chemical chemical unit having one or more reactive groups capable of forming a formation of a connection between the chemical entity and the nascent react with a reactive group also appearing on the original scaffold or a ន

naturally occurring or an artificial substance. Usually, a synthetic molecule is encoded molecule may be mediated by a bridging molecule. As an example, amine group a connection between these can be mediated by a dicarboxylic if the nascent encoded molecule and the chemical entity both comprise an acid. A synthetic molecule is in general produced in vitro and may be a not produced using the naturally translation system in an in vitro process. 22 8

eliminations of the encoded molecule may be attached to a building block The chemical entities that are precursors for structural additions or

WO 2005/026387

PCT/DK2004/000630

generally comprises an anti-codon. In some embodiments the building blocks prior to the participation in the formation of the reaction product leading the also comprise an affinity region providing for affinity towards the nascent final encoded molecule. Besides the chemical entity, the building block

molecule by a building block, which further comprises an anticodon. The anti-Thus, the chemical entities are suitably mediated to the nascent encoded codon serves the function of transferring the genetic information of the

- The chemical entities are preferably reacted without enzymatic interaction in some aspects of the invention. Notably, the reaction of the chemical entities is preferably not mediated by ribosomes or enzymes having similar activity. transfer of genetic information and chemical entity may occur in any order. building block in conjunction with the transfer of a chemical entity. The 9
 - In other aspects of the invention, enzymes are used to mediate the reaction between a chemical entity and a nascent encoded molecule. 5

According to certain aspects of the invention the genetic information of the anti-codon is transferred by specific hybridisation to a codon on a nucleic

- genetic information of the anti-codon to the nascent complex by an extension acid template. Another method for transferring the genetic information of the complex, e.g. by ligation. A still further method involves transferring the complementary to the anti-codon and attach this oligonucleotide to the anti-codon to the nascent complex is to anneal an oligonucleotide. ន
 - reaction using a polymerase and a mixture of dNTPs. 32

The chemical entity of the building block may in most cases be regarded as a molecule. In other cases the chemical entity provides for the eliminations of precursor for the structural entity eventually incorporated into the encoded

to a nascent encoded molecule it is to be understood that not necessarily all present application with claims is stated that a chemical entity is transferred chemical units of the nascent encoded molecule. Therefore, when it in the ഉ

the atoms of the original chemical entity is to be found in the eventually formed encoded molecule. Also, as a consequence of the reactions involved in the connection, the structure of the chemical entity can be changed when it appears on the nascent encoded molecule. Especially, the cleavage resulting in the release of the entity may generate a reactive group which in a subsequent step can participate in the formation of a connection between a nascent complex and a chemical entity.

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The chemical entity of the building block comprises at least one reactive

group capable of participating in a reaction which results in a connection between the chemical entity of the building block and another chemical entity or
a scaffold associated with the nascent complex. The number of reactive
groups which appear on the chemical entity is suitably one to ten. A building
block featuring only one reactive group is used i.a. in the end positions of

are suitable for the formation of the body part of a polymer or scaffolds capable of being reacted further. One, two or more reactive groups intended for
the formation of connections, are typically present on scaffolds. Non-limiting
examples of scaffolds are opiates, steroids, benzodiazepines, hydantoines,
and peptidylphosphonates.

The reactive group of the chemical entity may be capable of forming a direct connection to a reactive group of the nascent complex or the reactive group of the building block may be capable of forming a connection to a reactive group of the nascent complex through a bridging fill-in group. It is to be understood that not all the atoms of a reactive group are necessarily maintained in the connection formed. Rather, the reactive groups are to be regarded as precursors for the structure of the connection.

30 The subsequent cleavage step to release the chemical entity from the building block can be performed in any appropriate way. In an aspect of the invention the cleavage involves usage of a chemical reagent or an enzyme. The

WO 2005/026387

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PCT/DK2004/000630

cleavage results in a transfer of the chemical entity to the nascent encoded molecule or in a transfer of the nascent encoded molecule to the chemical entity of the building block. In some cases it may be advantageous to introduce new chemical groups as a consequence of linker cleavage. The new

5 chemical groups may be used for further reaction in a subsequent cycle, elther directly or after having been activated. In other cases it is desirable that no trace of the linker remains after the cleavage.

In another aspect, the connection and the cleavage is conducted as a simul-

taneous reaction, i.e. either the chemical entity of the building block or the nascent encoded molecule is a leaving group of the reaction. In some aspects of the invention, it is appropriate to design the system such that the connection and the cleavage occur simultaneously because this will reduce the number of steps and the complexity. The simultaneous connection and

15 cleavage can also be designed such that either no trace of the linker remains or such that a new chemical group for further reaction is introduced, as described above

The attachment of the chemical entity to the building block, optionally via a suitable spacer can be at any entity available for attachment, e.g. the chemical entity can be attached to a nucleobase or the backbone. In general, it is preferred to attach the chemical entity at the phosphor of the internucleoside linkage or at the nucleobase. When the nucleobase is used for attachment of the chemical entity, the attachment point is usually at the 7 position of the

purines or 7-deaza-purins or at the 5 position of pyrimidines. The nucleotide may be distanced from the reactive group of the chemical entity by a spacer moiety. The spacer may be designed such that the conformational spaced sampled by the reactive group is optimized for a reaction with the reactive group of the nascent encoded molecule.

The encoded molecules may have any chemical structure. In a preferred aspect, the encoded molecule can be any compound that may be

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synthetic molecule is a scaffolded molecule. The term "encoded molecule" synthesized in a component-by-component fashion. In some aspects the synthetic molecule is a linear or branched polymer. In another aspect the also comprises naturally occurring molecules like a-polypeptides etc, however produced in vitro usually in the absence of enzymes, like ribosomes. In certain aspects, the synthetic molecule of the library is a non-α-വ

The encoded molecule may have any molecular weight. However, in order to be orally available, it is in this case preferred that the synthetic molecule has a molecular weight less than 2000 Daltons, preferably less than 1000 Dalton, and more preferred less than 500 Daltons. 9

some aspects, the library comprises 1,000 or more different complexes, more more than two different complexes are desired to obtain a higher diversity. In preferred 1,000,000 or more different complexes. The upper limit for the size The size of the library may vary considerably pending on the expected result of the inventive method. In some aspects, it may be sufficient that the library of the library is only restricted by the size of the vessel in which the library is comprised. It may be calculated that a vial may comprise up to 10¹⁴ different comprises two, three, or four different complexes. However, in most events, 5

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Extension reaction

- recognise the double helix as a substrate. After binding of the polymerase to sufficient complementary to an identifier oligonucleotide for a polymerase to The extension reaction requires a primer, a polymerase as well as a collecproceed. An extension product may be obtained in the event the primer is lion of deoxyribonucleotide triphosphates (abbreviated dNTP's herein) to the double helix, the deoxyribonucleotide triphosphates (blend of dATP, ဓ 22
- dCTP, dGTP, and dTTP) are incorporated into the extension product using the identifier oligonucleotide as template. The conditions allowing for the exten-

WO 2005/026387

PCT/DK2004/000630

merase and the mixture of dNTP's are generally included in a buffer which is merase has a sufficient activity. To facilitate the extension process the polysion reaction to occur usually includes a suitable buffer. The buffer may be any aqueous or organic solvent or mixture of solvents in which the poly-

- process comprises the following: 50 mM KCl; 10 mM Tris-HCl at pH 8.3; 1.5 added to the identifier oligonucleotide and primer mixture. An exemplary kit mM MgCl2; 0.001% (wt/vol) gelatin, 200 µM dATP; 200 µM dTTP; 200 µM comprising the polymerase and the nNTP's for performing the extension dCTP; 200 µM dGTP; and 2.5 units Thermus aquaticus (Taq) DNA poly-
- merase I (U.S. Pat. No. 4,889,818) per 100 microliters (µI) of buffer. 9

The primer may be selected to be complementary to one or more codons or length of the codons, however, the primers usually are at least about 11 nuparts of such codons. The length of the primers may be determined by the

- for an efficient extension by the polymerase. The presence or absence of one cleotides in length, more preferred at least 15 nucleotides in length to allow or more codons is indicated by the presence of or absence of an extension product. The extension product may be measured by any suitable method, such as size fractioning on an agarose gel and stalning with ethidium bro-5
 - mide. 2

sion product. The thermocycling is typically carried out by repeatedly increasprimer is thermocycled to obtain a sufficient number of copies of the exten-In a preferred embodiment the admixture of identifier oligonucleotide and

- decreasing can be continuous, but is preferably phasic with time perlods of relative temperature stability at each of temperatures favouring polynucleorange whose lower limit is about 30 degrees Celsius (30°C) to about 55°C and whose upper limit is about 90°C to about 100° C. The increasing and ing and decreasing the temperature of the mixture within a temperature 23
 - tide synthesis, denaturation and hybridization. ဓ္တ

the result may be used to verify the presence or absence of a specific chemi-When a single complex is analysed in accordance with the present method, cal entity during the formation of the display molecule. The formation of an extension product is indicative of the presence of an oligonucleotide part

complementary to the primer in the identifier oligonucleotide. Conversely, the Selecting the sequence of the primer such that it is complementary to one or absence of an extension product is indicative of the absence of an oligonucleotide part complementary to the primer in the identifier oligonucleotide. more codons will therefore provide information of the structure of the encoded molecule coded for by this codon(s). S 9

cleotide and the primer oligonucleotide, a second primer complementary to a In a preferred aspect of the invention, in the mixture of the identifier oligonusequence of the extension product is included. The second primer is also

primer is well known to skilled person in the art and is generally referred to as termed reverse primer and ensures an exponential increase of the number of claims. In one embodiment of the invention the reverse primer is annealed to polymerase chain reaction (abbreviated PCR) in the present application with a part of the extension product downstream, i.e. near the 3'end of the extenneals to an upstream position of the identifier oligonucleotide, preferably begonucleotide. In another embodiment, the first primer (forward primer) anfore the coding part, and the reverse primer anneals to a sequence of the sion product, or a part complementing the coding part of the identifier oilproduced extension products. The method using a forward and reverse extension product complementing one or more codons or parts thereof. 5 ឧ 22

subsequently stained with ethidium bromide. Under UV illumination bands of amplicons becomes visible. It is possible to incorporate the staining agent in The amplicons resulting from the PCR process may be stained during or folan example, amplicons from the PCR process is run on an agarose gel and lowing the reaction to ease the detection. A staining after the PCR process may be prepared with e.g. ethidium bromide or a similar staining agent. As ဓ္တ

WO 2005/026387

PCT/DK2004/000630

the agarose gel or to allow a solution of the staining agent to migrate through the amplification proceeds it will incorporate in the double helix. The intercathe gel. The amplicons may also be stained during the PCR process by an intercalating agent, like CYBR. In presence of the intercalating agent while ation agent may then be made visible by irradiation by a suitable source.

cific amplicon. Thus, it is possible to quantify the occurrence of a codon in an The intensity of the staining is informative of the relative abundance of a speidentifier oligonucleotide. When a library of bifunctional complexes has been

- which has been selected can be quantified using this method. As an example PCR amplifications with different primers in separate compartments and the subjected to a selection the codons in the pool of identifier oligonucleotides a sample of the selected identifier oligonucleotides is subjected to various PCR product of each compartment is analysed by electrophoresis in the 2
- presence of ethidium bromide. The bands that appear can be quantified by a densitometric analysis after irradiation by ultraviolet light and the relative abundance of the codons can be measured. 5

when biotin is used as label and anti-digoxigenin when digoxigenin is used as biotin or digoxigenin. A PCR-ELISA analysis may subsequently be performed Alternatively, the primers may be labelled with a suitable small molecule, like ncludes the application of a solid support covered with streptavidin or avidin based on the amplicons comprising the small molecule. A preferred method the label. Once captured, the amplicons can be detected using an enzyme-2

labelled avidin or anti-dixigenin reporter molecule similar to a standard ELISA 22

ess "real time". Several real time PCR processes has been developed and all can be used in the evaluating step of the present invention and are include in cons, it is in a certain embodiment preferred to measure the extension proc-To avoid laborious post-PCR handling steps required to evaluate the amplithe suitable real time PCR process available to the skilled person in the art 9

PCT/DK2004/000630

. 20. the present scope of protection. The PCR reactions discussed below are of particular interest.

The monitoring of accumulating amplicons in real time has been made possi-

- ble by labelling of primers, probes, or amplicons with fluorogenic molecules.

 The real time PCR amplification is usually performed with a speed faster than the conventional PCR, mainly due to reduced cycles time and the use of sensitive methods for detection of emissions from the fluorogenic labels. The most commonly used fluorogenic oligoprobes rely upon fluorescent reso-
 - 10 nance energy transfer (FRET) between fluorogenic labels or between one flourophor and a dark or "black-hole" non-fluorescent quencher (NFQ), which disperse energy as heat rather than fluorescence. FRET is a spectroscopic process by which energy is passed between molecules separated by 10-100 A that have overlapping emission and absorption spectra. An advantage of many real time PCR methods is that they can be carried out in a closed system, i.e. a system which does not need to be opened to examine the result of the PCR. A closed system implies a reduced result turnaround, minimisation of the potential for carry-over contamination and the ability to closely scrutinise the essay's performance.

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The present real time PCR methods currently available to the skilled person can be classified into either amplicon sequence specific or non-specific methods. The basis for the non-specific detection methods is a DNA-binding fluorogenic molecule. Included in this class are the earliest and simplest approaches to real time PCR. Ethidium bromide, YO-PRO-1, and SYBR® green 1 all fluorescence when associated with double stranded DNA which is exposed to a suitable wavelength of light. This approach requires the fluorescent agent to be present during the PCR process and provides for a real time detection of the fluorescent agent as it is incorporated into the double

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stranded helix

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WO 2005/026387

PCT/DK2004/000630

The amplicons sequence specific methods includes, but are not limited to, the TaqMan®, hairpin, LightCycler®, Sunrise®, and Scorpion® methods. The LightCycler® method also designated "HybProbes" make use of a pair of adjacent, fluorogenic hybridisation oligonucleotide probes. A first, usually the

- usually the downstream probe is commonly labelled with either a Light cycler Red 640 or Red 705 acceptor fluorophore a the 5' terminus so that when both oligoprobes are hybridised the two fluorophores are located in close proximity, such as within 10 nm, of each other. The close proximity provides
- 10 for the emission of a fluorescence when Irradiated with a suitable light source, such a blue dlode in case of the LightCycler[®]. The ragion for annealing of the probes may be any suitable position that does not interfere with the primer annealing. In a suitable setup, the site for binding the probes are positioned downstream of the codon region on the identifier oligonucleotide. Alternatively, when a reverse primer is used, the region for annealing the
- ternatively, when a reverse primer is used, the region for annealing the probes may be at the 3' end of the strand complementing the identifier oll-gonucleotide. Another embodiment of the LightCycler method includes that the pair of oligonucleotide probes are annealed to one or more codons and primer sites extenior to the coding part of the identifier oligonucleotide are used for PCR amplification.

The TaqMan® method, also referred to as the 5' nuclease or hydrolysis method, requires an oligoprobe, which is attached to a reporter flourophor, such as 6-carboxy-fluoroscein, and a quencher fluorophore, such as 6-

- carboxy-tetramethyl-rhodamine, at each end. When In close proximity, I.e. annealed to an identifier oligonucleotide, or a sequence complementing the identifier oligonucleotide, the quencher will "hijack" the emissions that have resulted from the excitation of the reporter. As the polymerase progresses along the relevant strand, it displaces and the hydrolyses the oligoprobe via
- 30 its 5'--3' endonuclease activity. Once the reporter is removed from the extinguishing influence of the quencher, it is able to release excitation energy at a wavelength that can be monitored by a suitable instrument, such as ABI

Prism® 7700. The fractional cycle number at which the real-time fluorescence signal mirrors progression of the reaction above the background noise is normally used as an indicator of successful identifier oligonucleotide amplification. This threshold cycle (C_T) is defined as the PCR cycle in which the

- standard deviations of the mean base line fluorescence. The C_T is proportional to the number of identifier oligonucleotide copies present in the sample.

 The TaqMan probe is usually designed to hybridise at a position downstream of a primer binding site, be it a forward or a reverse primer. When the primer is designed to an everse primer of a primer binding site, and a primer of a primer binding site, and a primer or a primer of a primer binding site, and a primer or a primer or a primer binding site, and a primer or a primer or a primer binding site, and a primer or a primer or a primer binding site, and a primer or a primer or a primer binding site, and a primer or a primer or a primer binding site, and a primer or a primer or a primer binding site, and a primer or a primer binding site, and a primer or a primer binding site, and a primer binding site of a primer binding site, and a primer binding site of a primer
- 10 is designed to anneal to one or more codons of the identifier oligonucleotide, the presence of these one or more codons is indicated by the emittance of light. Furthermore, the quantity of the identifier oligonucleotides comprising the one or more codons may be measured by the $C_{\rm T}$ value.
- The Hairpin method involves an oligoprobe, in which a fluorophore and a quencher are positioned at the termini. The labels are hold in close proximity by distal stem regions of homologous base pairing deliberately designed to create a hairpin structure which result in quenching either by FRET or a direct energy transfer by a collisional mechanism due to the intimate proximity of the labels. When direct energy transfer by a collision mechanism is used
- the quencher is usually different from the FRET mechanism, and is suitably 4-(4'-dimethylamino-phenylazo)-benzene (DABCYL). In the presence of a complementary sequence, usually downstream of a primer, or within the bounds of the primer binding sides in case of more than one a single primer, the oligoprobe will hybridise, shifting into an open configuration. The fluorophore is now spatially removed from the quencher's influence and fluorescence emissions are monitored during each cycle. In a certain aspect, the hairpin probe may be designed to anneal to a codon in order to detect this
- 30 suitable if codons only differs from each other with a single or a few nucleo-tides, because is in well-known that the occurrence of a mismatch between a hairpin oligoprobe and its target sequence has a greater destabilising effect

codon if present on the identifier oligonucleotide. This embodiment may be

WO 2005/026387 PCT/DK2004/000630

23

on the duplex than the introduction of an equivalent mismatch between the target oligonucleotide and a linear oligoprobe. This is probably because the hairpin structure provides a highly stable alternate conformation.

- 5 The Sunrise and Scorpion methods are similar in concept to the hairpin oligoprobe, except that the label becomes irreversible incorporated In to the PCR product. The Sunrise method involves a primer (commercially available as AmplifluorTM hairpin primers) comprising a 5' fluorophore and a quencher, e.g. DABCYL. The labels are separated by complementary stretches of seeg.
- 10 quence that create a stem when the sunrise primer is closed. At the 3' terminus is a target specific primer sequence. In a preferred embodiment the target sequence is a codon, optionally more codons. The sunrise primer's sequence is intended to be duplicated by the nascent complementary stand and, in this way, the stem is destabilised, the two fluorophores are held apart,
- usually between 15 and 25 nucleotides, and the fluorophore is free to emit its excitation energy for monitoring. The Scorpion primer resembles the sunrise primer, but derivate in having a molety that blocks duplication on the signal-ling portion of the scorpion primer. The blocking moiety is typically hexethylene glycol. In addition to the difference in structure, the function of the scor-
- 20 pion primers differs slightly In that the 5' region of the oligonucleotide Is designed to hybridise to a complementary region within the amplicons. In a certain embodiment the complementary region is a codon on the identifier oligonucleotide. The hybridisation forces the labels apart disrupting the halrpin and permitting emission in the same way as the halrpin probes.

Methods for forming a library of complexes

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The complexes comprising an identifier part having two or more codons that code for reactants that have reacted in the formation of the encoded molecule part of the complex may be formed by a variety of processes. Generally,

30 the preferred methods can be used for the formation of virtually any kind of encode molecule. Suitable examples of processes include prior art methods disclosed in WO 93/20242, WO 93/06121, WO 00/23458, WO 02/074929,

and WO 02/103008, the content of which being incorporated herein by reference as well as methods of the present applicant not yet public available, including the methods disclosed in DK PA 2002 01955 filed 19 December 2002, and DK PA 2003 00430 filed 20 March 2003. Any of these methods may be used, and the entire content of the patent applications are included herein by reference.

Below four preferred embodiments are described. A first embodiment disclosed in more detail in WO 02/103008 is based on the use of a polymerase to incorporate unnatural nucleotides as building blocks. Initially, a plurality of template oligonucleotides is provided. Subsequently primers are

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plurality of template oligonucleotides is provided. Subsequently primers are annealed to each of the templates and a polymerase is extending the primer using nucleotide derivatives which have appended chemical entities.

Subsequent to or simultaneously with the incorporation of the nucleotide derivatives, the chemical entities are reacted to form a reaction product. The encoded molecule may be post-modified by cleaving some of the linking moieties to better present the encoded molecule.

Several possible reaction approaches for the chemical entities are apparent.

First, the nucleotide derivatives can be incorporated and the chemical entities subsequently polymerised. In the event the chemical entities each carry two reactive groups, the chemical entities can be attached to adjacent chemical entities by a reaction of these reactive groups. Exemplary of the reactive groups are amine and carboxylic acid, which upon reaction form an amide bond. Adjacent chemical entities can also be linked together using a linking

25 bond. Adjacent chemical entitles can also be linked together using a linking or bridging moiety. Exemplary of this approach is the linking of two chemical entities each bearing an amine group by a bi-carboxylic acid. Yet another approach is the use of a reactive group between a chemical entity and the nucleotide building block, such as an ester or a thioester group. An adjacent building block having a reactive group such as an amine may cleave the interspaced reactive group to obtain a linkage to the chemical entity, e.g. by an amide linking group.

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PCT/DK2004/000630

A second embodiment for obtainment of complexes pertains to the use of hybridisation of building blocks to an Identifier oligonucleotide and reaction of chemical entities attached to the building blocks in order to obtain a reaction product. This approach comprises that templates are contacted with a

product. This approach comprises that templates are contacted with a plurality of building blocks, wherein each building block comprises an anti-codon and a chemical entity. The anti-codons are designed such that they recognise a sequence, I.e. a codon, on the template. Subsequent to the annealing of the anti-codon and the codon to each other a reaction of the chemical entity is effected.

The template may be associated with a scaffold. Building blocks bringing chemical entities in may be added sequentially or simultaneously and a reaction of the reactive group of the chemical entity may be effected at any

time after the annealing of the building blocks to the template.

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A third embodiment for the generation of a complex includes chemical or enzymatical ligation of building blocks when these are lined up on a template. Initially, templates are provided, each having one or more codons. The

templates are contacted with building blocks comprising anti-codons linked to chemical entities. The two or more anti-codons annealed on a template are subsequently ligated to each other and a reaction of the chemical entities is effected to obtain a reaction product. The method is disclosed in more detail in DK PA 2003 00430 filed 20 March 2003.

A fourth embodiment makes use of the extension by a polymerase of an affinity sequence of the nascent complex to transfer the anti-codon of a building block to the nascent complex. The method implies that a nascent complex comprising a scaffold and an affinity region is annealed to a building block comprising a region complementary to the affinity section.

33

block comprising a region complementary to the affinlty section.

Subsequently the anti-codon region of the building block is transferred to the nascent complex by a polymerase. The transfer of the chemical entity may

the anti-codon. This method is disclosed in detail in DK PA 2002 01955 filed be transferred prior to, simultaneously with or subsequent to the transfer of 19 December 2002. After or simultaneously with the formation of the reaction product some of the linkers to the template may be cleaved, however at least one linker must be maintained to provide for the complex. S

sequence of nucleotides, i.e. an oligonucleotide. Each nucleotide monomer is bone. The back bone may in some cases be subdivided into a sugar moiety normally composed of two parts, namely a nucleobase moiety, and a back-The nucleotides used in the present invention may be linked together in a and an internucleoside linker. 9

The nucleobase moiety may be selected among naturally occurring nucleobases as well as non-naturally occurring nucleobases. Thus, "nucleobase" heterocyclic analogues and tautomers thereof. Illustrative examples of nuincludes not only the known purine and pyrimidine hetero-cycles, but also

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hydroxy-5-methyl-4-triazolopyridine, isocytosine, isoguanine, inosine and the bases are adenine, guanine, thymine, cytosine, 5-methylcytosine, and uracil, N*,N*-ethanocytosin, N⁶,N⁸-ethano-2,6-diamino-purine, 5-methylcytosine, 5-(C3-C3)-alkynylcytosine, 5-fluorouracil, 5-bromouracil, pseudoisocytosine, 2diaminopurine, 8-oxo-N⁸-methyladenine, 7-deazaxanthine, 7-deazaguanine, cleobases are adenine, guanine, thymine, cytosine, uracil, purine, xanthine, No. 5,432,272. The term "nucleobase" is intended to cover these examples "non-naturally occurring" nucleobases described in Benner et al., U.S. Pat as well as analogues and tautomers thereof. Especially interesting nucleowhich are considered as the naturally occurring nucleobases in relation to therapeutic and diagnostic application in humans. 22 ನ ജ

Examples of suitable specific pairs of nucleobases are shown below:

WO 2005/026387

PCT/DK2004/000630

27

ynthetic Base Patr

Suitable examples of backbone units are shown below (B denotes a nucleo-

propriate part of an PNA or a six-member ring. Suitable examples of possible and 2'4'-O-methylene-ribose (LNA). Suitably the nucleobase is attached to pentoses include ribose, 2'-deoxyribose, 2'-O-methyl-ribose, 2'-flour-ribose, The sugar moiety of the backbone is suitably a pentose but may be the apthe 1' position of the pentose entity.

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end of a succeeding monomer when the sugar moiety of the backbone is a phosphoramidate, phosphotriester, and phosphodithioate. Furthermore, the An internucleoside linker connects the 3' end of preceding monomer to a 5' pentose, like ribose or 2-deoxyribose. The internucleoside linkage may be the natural occurring phospodiester linkage or a derivative thereof. Examples of such derivatives include phosphorothioate, methylphosphonate, 9

WO 2005/026387

PCT/DK2004/000630

internucleoside linker can be any of a number of non-phosphorous-containing inkers known in the art.

and inosine. Inosine is a non-specific pairing nucleoside and may be used as forming part of the DNA as well as the RNA family connected through phosmembers of the RNA family include adenosine, guanosine, uridine, cytidine, Preferred nucleic acid monomers include naturally occurring nucleosides oxyadenosine, deoxyguanosine, deoxythymidine, and deoxycytidine. The phodiester linkages. The members of the DNA family Include de-2

universal base because inosine can pair nearly isoenergetically with A, T, and C. Other compounds having the same ability of non-specifically basepairing with natural nucleobases have been formed. Suitable compounds which may be utilized in the present invention includes among others the compounds depicted below 9

Examples of Universal Bases:

3-Nitropyrrole N*-8aza-7deazaadenine

5-NitroIndole

Inosine

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Nebularine

Building block

The chemical entities that are precursors for structural additions or elimina-

tions of the encoded molecule may be attached to a building block prior to S

WO 2005/026387

PCT/DK2004/000630

the participation in the formation of the reaction product leading the final encoded molecule. Besides the chemical entity, the building block generally comprises an anti-codon.

tween the chemical entity of the building block and another chemical entity or a scaffold associated with the nascent complex. The connection is facilitated group capable of participating in a reaction which results in a connection befolds capable of being reacted further. One, two or more reactive groups inbuilding block featuring only one reactive group is used i.a. in the end posilions of polymers or scaffolds, whereas building blocks having two reactive by one or more reactive groups of the chemical entity. The number of reacgroups are suitable for the formation of the body part of a polymer or scaftended for the formation of connections, are typically present on scaffolds. The chemical entity of the building block comprises at least one reactive tive groups which appear on the chemical entity is suitably one to ten. A ťΩ 9

connection to a reactive group of the nascent complex or the reactive group The reactive group of the building block may be capable of forming a direct of the building block may be capable of forming a connection to a reactive

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- derstood that not all the atoms of a reactive group are necessarily maintained in the connection formed. Rather, the reactive groups are to be regarded as group of the nascent complex through a bridging fill-in group. It is to be unprecursors for the structure of the connection. 2
- ng block can be performed in any appropriate way. In an aspect of the Invenor in a transfer of the nascent encoded molecule to the chemical entity of the The subsequent cleavage step to release the chemical entity from the buildresults in a transfer of the chemical entity to the nascent encoded molecule tion the cleavage involves usage of a reagent or an enzyme. The cleavage . 22
- groups may be used for further reaction in a subsequent cycle, either directly chemical groups as a consequence of linker cleavage. The new chemical building block. In some cases it may be advantageous to introduce new 8

32

or after having been activated. In other cases it is desirable that no trace of the linker remains after the cleavage.

complexity. The simultaneous connection and cleavage can also be designed nascent encoded molecule is a leaving group of the reaction. In general, it is In another aspect, the connection and the cleavage is conducted as a simultaneous reaction, i.e. either the chemical entity of the building block or the occur simultaneously because this will reduce the number of steps and the preferred to design the system such that the connection and the cleavage such that either no trace of the linker remains or such that a new chemical group for further reaction is introduced, as described above. Ŋ 9

inkage or at the nucleobase. When the nucleobase is used for attachment of preferred to attach the chemical entity at the phosphor of the internucleoside suitable spacer can be at any entity available for attachment, e.g. the chemipurines or 7-deaza-purins or at the 5 position of pyrimidines. The nucleotide may be distanced from the reactive group of the chemical entity by a spacer cal entity can be attached to a nucleobase or the backbone. In general, it is The attachment of the chemical entity to the building block, optionally via a the chemical entity, the attachment point is usually at the 7 position of the sampled by the reactive group is optimized for a reaction with the reactive molety. The spacer may be designed such that the conformational space group of the nascent encoded molecule or reactive site. 5

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The anticodon complements the codon of the identifier sequence and generally comprises the same number of nucleotides as the codon. The anticodon may be adjoined with a fixed sequence, such as a sequence complementing a framing sequence. 22

Various specific building blocks are envisaged. Building blocks of particular interest are shown below. 9

WO 2005/026387

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PCT/DK2004/000630

Specific Building blocks

Building blocks transferring a chemical entity to a reciplent nucleophilic group

The building block indicated below is capable of transferring a chemical entity

lower horizontal line illustrates the building block and the vertical line illus-(CE) to a recipient nucleophilic group, typically an amine group. The bold

ring serves as an activator, i.e. a labile bond is formed between the oxygen trates a spacer. The 5-membered substituted N-hydroxysuccinimid (NHS) atom connected to the NHS ring and the chemical entity. The labile bond

may be cleaved by a nucleophilic group, e.g. positioned on a scaffold

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activator, i.e. a labile bond is formed between the oxygen atom connected to The 5-membered substituted N-hydroxysuccinImid (NHS) ring serves as an

- ing group of the reaction. When the chemical entity is connected to the actinucleophilic group, e.g. positioned on a scaffold, to transfer the chemical entity to the scaffold, thus converting the remainder of the fragment into a leavthe NHS ring and the chemical entity. The labile bond may be cleaved by a ator through an carbonyl group and the recipient group is an amine, the 5
- provisional patent application No. 60/434,439, the content of which are incorbond formed on the scaffold will an amide bond. The above building block is he subject of the Danish patent application No. PA 2002 01946 and the US porated herein in their entirety by reference. 8
- Another building block which may form an amide bond is 22

R may be absent or NO2, CF3, halogen, preferably Cl, Br, or I, and Z may be 2002 with the title "A building block capable of transferring a functional entity No. PA 2002 0951 and US provisional patent application filed 20 December S or O. This type of building block is disclosed in Danish patent application to a recipient reactive group". The content of both patent application are incorporated herein in their entirety by reference.

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10 · group thereby transferring the chemical entity -(C=O)-CE' to said nucleo-A nucleophilic group can cleave the linkage between Z and the carbonyl philic group. Building blocks transferring a chemical entity to a recipient reactive group forming a C=C bond

recipient aldehylde group thereby forming a double bond between the carbon A building block as shown below are able to transfer the chemical entity to a of the aldehyde and the chemical entity 5

WO 2005/026387

35

PCT/DK2004/000630

content of both patent applications are incorporated herein in their entirety by The above building block is comprised by the Danish patent application No. tional entity to a recipient reactive group forming a C=C double bond". The cember 2002 with the title "A building block capable of transferring a func-DK PA 2002 01952 and the US provisional patent application filed 20 De-

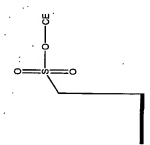
Building blocks transferring a chemical entity to a recipient reactive group forming a C-C bond 5

The below building block is able to transfer the chemical entity to a recipient group thereby forming a single bond between the receiving molety, e.g. a scaffold, and the chemical entity.

DK PA 2002 01947 and the US provisional patent application No 60/434,428. The above building block is comprised by the Danish patent application No. The content of both patent applications are incorporated herein in their en-

tirety by reference. 8

Another building block capable of transferring a chemical entity to a receiving reactive group forming a single bond is



The receiving group may be a nucleophile, such as a group comprising a hetero atom, thereby forming a single bond between the chemical entity and the hetero atom, or the receiving group may be an electronegative carbon

10 atom, thereby forming a C-C bond between the chemical entity and the scaffold.

The chemical entity attached to any of the above building blocks may be a selected from a large arsenal of chemical structures. Examples of chemical to entities are

H or entities selected among the group consisting of a C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkenyl, C₂-C₆ alkanyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, and heteroaryl, said group being substituted with 0-3 R⁴, 0-3 R⁵ and 0-3 R⁹ or C₁-C₃ alkylene-NR², C₁-C₃ alkylene-NR⁴C(O)R⁸, C₁-C₃ al-

20 kylene-NR⁴C(O)OR⁸, C₁-C₂ alkylehe-O-NR²s, C₁-C₂ alkylene-O-NR⁴C(O)R⁸, C₁-C₂ alkylene-O-NR⁴C(O)OR⁸ substituted with 0-3 R⁹.

Where R⁴ is H or selected independently among the group consisting

2

of C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl, heteroaryl, said group being substituted with 0-3 R⁹ and

25 R^5 is selected independently from -N₃, -CNO, -C(NOH)NH₂, -NHOH, -NHNHR⁶, -C(O)R⁶, -SnR⁸3, -B(OR⁶)₂, -P(O)(OR⁶)₂ or the group consisting of

WO 2005/026387

PCT/DK2004/000630

 C_2 - C_6 alkenyl, C_2 - C_6 alkynyl, C_4 - C_8 alkadienyl said group being substituted with 0-2 R 7 ,

37

where R⁶ is selected independently from H, C₁-C₈ alkyl, C₃-C₇ cycloalkyl, aryl or C₁-C₈ alkylene-aryl substituted with 0-5 halogen atoms selected from -F, -Cl, -Br, and -l; and R⁷ is independently selected from -NO₂,

-COOR⁶, -COR⁹, -CN, -OSIR⁶s, -OR⁸ and -NR⁵.

R⁸ is H, C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₃-C₇ cycloalkyl, aryl or
C₁-C₆ alkylene-aryl substituted with 0-3 substituents independently selected from -F, -Cl, -NO₂, -R³, -OR³, -SiR³,

10 R³ is =O, -F, -Cl, -Br, -I, -CN, -NO₂, -OR⁶, -NR⁶, -NR⁶-C(O)R⁶, -S(O)R⁶, -S(O)₂R⁶, -COOR⁶, -C(O)NR⁵₂ and -S(O)₂NR⁶, -S(O)₂N

Cross-link cleavage building blocks

15 It may be advantageous to split the transfer of a chemical entity to a recipient reactive group into two separate steps, namely a cross-linking step and a cleavage step because each step can be optimized. A suitable building block for this two step process is illustrated below:

Initally, a reactive group appearing on the functional entity precursor (abbreviated FEP) reacts with a recipient reactive group, e.g. a reactive group appearing on a scaffold, thereby forming a cross-link. Subsequently, a cleavage

is performed, usually by adding an aqueous oxidising agent such as 12, Br2,

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Cl₂, H*, or a Lewis acid. The cleavage results in a transfer of the group HZ-FEP- to the recipient moiety, such as a scaffold.

In the above formula

Z is O, S, NR⁴

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Q is N, CR1

C1-60-alkylene, C1-6S-alkylene, NR1-alkylene, C1-6alkylene-O, C1-6alkylene-S P is a valence bond, O, S, NR 4 , or a group C $_{57}$ arylene, C $_{16}$ allkylene, option said group being substituted with 0-3 R⁴, 0-3 R⁵ and 0-3 R⁹ or C₁-C₃ alkylene-NR², C₁-C₃ alkylene-NR⁴C(O)R⁸, C₁-C₃ alkylene-NR⁴C(O)OR⁸

C₁-C₂ alkylene-O-NR⁴2, C₁-C₂ alkylene-O-NR⁴C(O)R⁸, C₁-C₂ alkylene-O-NR⁴C(O)OR⁸ substituted with 0-3 R⁹,

은

B is a group comprising D-E-F, in which

D is a valence bond or a group C1-6alkylene, C1-6alkenylene, C1.

ealkynylene, Cs-7arylene, or Cs-7heteroarylene, said group optionally being substituted with 1 to 4 group R11, 5

ealkylene, C₁₋₆alkenylene, C₁₋₆alkynylene, C₅₋₇arylene, or C₅₋₇heteroarylene, E is, when present, a valence bond, O, S, NR⁴, or a group C₁. said group optionally being substituted with 1 to 4 group R¹¹

F is, when present, a valence bond, O, S, or NR4

8.

A is a spacing group distancing the chemical structure from the complementing element, which may be a nucleic acid,

kylene-NR⁴2, C₁-C₃ alkylene-NR⁴C(O)R⁸, C₁-C₃ alkylene-NR⁴C(O)OR⁸, C₁-C₂ group consisting of H, C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₄-C₆ alkadi-Ri, R2, and R3 are independent of each other selected among the enyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 R⁴, 0-3 R⁵ and 0-3 R⁹ or C₁-C₃ alalkylene-O-NR⁴2, C₁-C₂ alkylene-O-NR⁴C(O)R⁸, C₁-C₂ alkylene-O-NR⁴C(O)OR⁸ substituted with 0-3 R⁹, 23

heteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 R4, 0-3 FEP is a group selected among the group consisting of H, C₁-C₆ alkyl, C2-C6 alkenyl, C2-C6 alkynyl, C4-C8 alkadienyl, C3-C7 cycloalkyl, C3-C7 cyclo-ဓ

WO 2005/026387

PCT/DK2004/000630

R⁵ and 0-3 R⁹ or C₁-C₃ alkylene-NR⁴2, C₁-C₃ alkylene-NR⁴C(O)R⁹, C₁-C₃ al-

kylene-NR⁴C(O)OR⁸, C₁-C₂ alkylene-O-NR⁴2, C₁-C₂ alkylene-O-NR⁴C(O)R⁸, C₁-C₂ alkylene-O-NR⁴C(O)OR⁸ substituted with 0-3 R⁹, of C₁-C₈ alkyl, C₂-C₆ alkenyl, C₂-C₈ alkynyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroalkyl, aryl, heteroaryl, said group being substituted with 0-3 R9 and

where R4 is.H or selected independently among the group consisting

-NHNHR 6 , -C(O)R 6 , -SnR 9 3, -B(OR 6)2, -P(O)(OR 9)2 or the group consisting of R⁵ is selected independently from -N₃, -CNO, -C(NOH)NH₂, -NHOH, Cz-Cs alkenyl, Cz-Cs alkynyl, C4-Cs alkadienyl said group being substituted with 0-2 R7, 2

where R⁶ is selected independently from H, C₁-C₆ alkyl, C₃-C₇ cycloalkyl, aryl or C₁-C₆ alkylene-aryl substituted with 0-5 halogen atoms selected from -F, -Cl, -Br, and \dashv ; and R^7 is independently selected from $\dashv \mathsf{NO}_2$, -COOR⁶, -COR⁶, -CN, -OSIR⁶₃, -OR⁶ and -NR⁶₂.

C₁-C₈ alkylene-aryl substituted with 0-3 substituents independently selected R⁸ is H, C₁-C₈ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₃-C₇ cycloalkyl, aryl or -NR $^{\rm e}$ -C(O)OR $^{\rm e}$, -S(O)R $^{\rm e}$, -S(O)2R $^{\rm e}$, -COOR $^{\rm e}$, -C(O)NR $^{\rm e}_2$ and R^9 is =0, -F, -CI, -Br, -I, -CN, -NO2, -OR 6 , -NR 2 2, -NR 6 -C(O)R 8 , from -F, -Cl, -NO₂, -R³, -OR³, -SiR³ 5

-S(O)₂NR⁶₂. 2

CH2, and R1, R2, and R3 is H. The bond between the carbonyl group and Z is In a preferred embodiment Z is O or S, P is a valence bond, Q is CH, B is cleavable with aqueous I2.

Partitioning

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property from the remainder of the library, may be referred to as the enrich-The partitioning step, by which the library of bifunctional molecules is subected to a condition partitioning one or more complexes having a certain

ment step or the selection step, as appropriate, and includes the screening of stics. Predetermined desirable characteristics can include binding to a target, the library for encoded molecules having predetermined desirable character-ജ

PCT/DK2004/000630

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ner which alters/modifies the target or the functional activity of the target, and catalytically changing the target, chemically reacting with a target in a mancovalently attaching to the target as in a suicide inhibitor.

- cording to the present invention is to enrich molecules with respect to binding involve mixing the library of complexes with the immobilized target of interest. affinity towards a target of interest. In a certain embodiment, the basic steps using either physical or physiological procedures. The method preferred ac-In theory, molecules of interest can be selected based on their properties Ŋ
 - without immobilisation of the target. Displayed molecules that bind to the target will be retained by a filter, size-exclusion chromatography etc, while nonimmobilization or by means of antibody binding or other high-affinity interac-The target can be attached to a column matrix or microtitre wells with direct tions. In another embodiment, the target and displayed molecules interact 15 9
 - separated by cleaving a physical connection to the encoded molecule or the entire complex may be eluted. It may be considered advantageously to per-... cleavage of the physical link between the synthetic molecule and the identifier, the identifier may be recovered from the media and optionally amplified form a chromatography step after or instead of the washing step. After the binding displayed molecules will be removed during a single or a series of wash steps. The identifiers of complexes bound to the target can then be
- before the decoding step. ೪
- A significant reduction in background binders may be obtained with increased ber of steps used in the washing procedure together with more stringent conditions will more efficiently remove non-binders and background binders. The specific binders. However, the washing step will also remove wanted binders washing volumes, repeating washing steps, higher detergent concentrations and prolonged incubation during washing. Thus, the more volume and numright stringency in the washing step can also be used to remove low-affinity 8 32

if too harsh conditions are used.

WO 2005/026387

PCT/DK2004/000630

tions should be as stringent as possible to remove background binding but to eins or other inert proteins and/or mild detergent such as Tween-20 and Triton X-100, may also be used to reduce the background. The washing condi-A blocking step, such as incubation of solid phase with skimmed milk pro-

vashing conditions are adjusted to maintain the desired affinity binders, e.g. retain specific binders that interact with the immobilized target. Generally, binders in the micromolar, nanomolar, or pocomolar range.

tor, inhibitor, drug, dye, nutrient, growth factor, cell, tissue, etc. without limita-The target can be any compound of interest. E.g. the target can be a protein, tigen, antibody, virus, substrate, metabolite, transition state analogue, cofacenzyme, renin, cyclooxygenase, 5-lipoxygenase, IIL- 1 0 converting enzyme, peptide, carbohydrate, polysaccharide, glycoprotein, hormone, receptor, antion. Suitable targets include, but are not limited to, anglotensin converting 9

- cytokine receptors, PDGF receptor, type II Inosine monophosphate dehydro-Abl/Her, phosphotases like PTP-1B, and fungal cytochrome P-450. Targets genase, B-lactamases, integrin, proteases like factor VIIa, kinases like Bcrcan include, but are not limited to, bradykinin, neutrophil elastase, the HIV proteins, including tat, rev, gag, int, RT, nucleocapsid etc., VEGF, bFGF, to
 - cells, glioblastomas, fibrin clots, PBMCs, hCG, lectins, selectins, cytokines, TGFB, KGF, PDGF, GPCR, thrombin, substance P, IgE, sPLA2, red blood ICP4, complement proteins, etc. 8

surface or a metal surface. The method of the invention may then be used to A target can also be a surface of a non-biological origin, such as a polymer dentify suitable coatings for such surfaces. 22

get without any interaction between the nucleic acid attached to the desirable In a preferred embodiment, the desIrable synthetic molecule acts on the tar-

arget aggregate can be partitioned from unbound complexes by a number of encoded molecule and the target. In one embodiment, the bound complexmethods. The methods include nitrocellulose filter binding, column chroma-ജ

tography, filtration, affinity chromatography, centrifugation, and other well known methods. A preferred method is size-exclusion chromatography. Briefly, the library of complexes is subjected to the target, which may include cules (e.g. encoded molecules which cross-react with other targets) may be used, will pass through the column. Additional undesirable encoded molecontact between the library and a column onto which the target is immobilised. Identifiers associated with undesirable encoded molecules, i.e. synthetic molecules not bound to the target under the stringency conditions

bodiment, the target is immobilized through a cleavable physical link, such as one more chemical bonds. The aggregate of the target and the complex may then be subjected to a size exclusion chromatography to separate the aggrethe column. The target may be immobilized in a number of ways. In one emremoved by counter-selection methods. Desirable complexes are bound to क 9

able linker, preferable orthogonal to a cleavable linker that attaches the target gate from the rest of the compounds in the media. The complex may then be temperature etc.). Alternatively, the complex may be provided with a cleaveluted from the target by changing the conditions (e.g., salt, pH, surfactant, to the solid support, at a position between the synthetic molecule and the

wards the targets. Just to mention a single type of orthogonal cleavable linkcan be cleaved by a chemical agent, and the linker separating the synthetic ages, one could attach the target to the solid support through a linkage that identifier. Subsequent to the size exclusion chromatography this cleavable linker is cleaved to separate the identifiers of complexes having affinity to-2 22

More specifically, the former linkage may be a disulphide bond that can be cleaved by a suitable reducing agent like DTT (dithiothreitol) and the latter molecule and the identifier may be selected as a photocleavable linkage. linkage may be a o-nitrophenyl group. There are other partitioning and screening processes which are compatible with this invention that are known to one of ordinary skill in the art. In one embodiment, the products can be fractionated by a number of common ജ

WO 2005/026387

PCT/DK2004/000630

methods and then each fraction is assayed for activity. The fractionization methods can include size, pH, hydrophobicity, etc. Inherent in the present method is the selection of encoded molecules on the

- selection), followed by positive selection with the desired target. As an examcules with a desired function and specificity. Specificity can be required durng the selection process by first extracting complexes which are capable of basis of a desired function; this can be extended to the selection of molenteracting with a non-desired "target" (negative selection, or counter-2
- eact to some extent with mammalian cytochrome P-450 (resulting in serious ected from a library by first removing those complexes capable of interacting side effects). Highly specific inhibitors of the fungal cytochrome could be seple, inhibitors of fungal cytochrome P-450 (fungicides) are known to crosswith the mammalian cytochrome, followed by retention of the remaining products which are capable of interacting with the fungal cytochrome. 9

Brief Description of the Figures

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probe) in the measurement of the presence or absence of a certain codon. Fig. 11 discloses two embodiments of using a Taqman probe (5' nuclease

Fig. 13 discloses the result of two experiments reported in example 3. Fig. 12 discloses a standard curve used in example 3. ೫

Example 3 - Illustrating the first aspect of the present invention

A preferred embodiment of the invention utilizing a universal Taqman probe

- along with flanking regions (light pattern). A universal Taqman probe anneals the universal PCR primers Pr.1 and Pr. 2. These primers could be the same to a region adjacent to the codon region, but within the amplicon defined by as used for amplification of the identifier oligonucleotides encoding binders is shown in Fig.11. Four codons are shown (P1 through P4; bold pattern) 22
 - Taqman probe annealing could be appended to the library identifier oligonuafter an enrichment process on a specific target. However, if minimal length templates are preferred during the encoding process, the region involved in ဗ္က

PCR primers. The Q-PCR reactions are preferably performed in a 96- or 384sponding to the region necessary for annealing of the Taqman probe would cleotides by e.g. overlap PCR, ligation, or by employing a long downstream PCR primer containing the necessary sequences. The added length correbe form 20 to 40 nts depending on the type of TaqMan probe and T_{A} of the well format on a real-time PCR thermocycling machine.

rO

position one. Similar primers are prepared for all codon sequences. For each codon sequence utilized to encode a specific BB in the library a Q-PCR reac-Fig. 11A shows the detection of abundance of a specific codon sequence in the PCR amplicon. The setup is most suited for cases where the codon contion is performed with a primer oligonucleotide complementary to the codon vided after the Taqman probe to provide for an exponential amplification of sequence in question. A downstream universal reverse primer Pr. 2 is prostitutes a length corresponding to a length suitable for a PCR primer.

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Fig. 11B shows the detection of abundance of a specific codon sequence in a region up- or downstream of the codon region which ensures extension of the specific codon position using a primer which is complementing a codon and a specific codon position. The number of specific primers and Q-PCR reactions sitions can be performed in a single run on four 96 wells micro titre plates (as primer in a PCR reaction only when annealed to the codon sequence in that the number of codon sequences times the number of codon positions. Thus, monitoring the abundance of 96 different codon sequences in 4 different poarchitecture allows for the decoding of a 8,5 x107 library of different encoded specific codon position in the library a Q-PCR reaction is performed with an needed to cover all codon sequences in all possible codon positions equals sequences. For each codon sequence utilized to encode a specific BB at a shown in Fig. 11B) or a single 384 well plate on a suitable instrument. This oligo complementary to the codon sequence in question as well as a short framing sequence. Similar primers are used for all the codons and framing 8 25 ဓ္က

PCT/DK2004/000630 WO 2005/026387

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with the two external PCR primers Pr.1 + Pr. 2. Theoretically, a similar rate of Quantification is performed relative to the amount of full-length PCR product obtained in a parallel control reaction on the same input material performed accumulation of this control amplicon compared to the accumulation of a

product utilizing a single codon + sequence specific primer would indicate a could be utilized. In theory, multiplex reactions employing up to 4 different Although the setups shown in Fig. 11A and 11B employ a Tagman probe strategy, other detection systems (SYBR green, Molecular Beacons etc.) 100% dominance of this particular sequence in the position in question. 2

fluorofors in the same reaction could increase throughput correspondingly,

centration. Three different chemical entitles are present in the first position of cules occurs is described in the following. Imagine that at the end of a selecthe encoded compounds, and each of these chemical entities are present in plates) are dominating the population and present at approx. the same contion scheme a pool of 3 ligand families (and the corresponding coding tem-An example of how a deconvolution process of a library of encoded mole-. 5

combination with one unique chemical entity out of 3 different chemical entiies in position P2. Only one chemical entity In position 3 gives rise to active binders, whereas any of a 20% subset of chemical entities (e.g. determined come of the initial codon profile analysis would be: 3 codon sequences are by charge, size or other characteristica) are present in position 4. The out-20

creased levels of 20% of the codon sequences (background levels of the reequally dominating in position P1, 3 other codon sequences in position P2, 1 to use an iterative Q-PCR ("IQ-PCR") strategy to perform a further deconvomaining 80% sequences) are seen in P4. In such cases it could be relevant unique codon sequence is dominant in P3 whereas somewhat similarly in-33

by taking the PCR products from the 3 individual wells that contained primers lution of a library after selection. Again with reference to the example above, giving the high yields in position P1, diluting the product appropriately and ဓ

performing a second round of Q-PCR on each of these identifier oligonucleotides separately, it would be possible to deduce which codon sequence(s) is preferred in P2 when a given codon sequence is present in P1.

Tagman MGB probe binding region: *=AATTCCAGCTTCTAGGAAGAC

P1 P2 P3 F

5' - CAGCTTGGACACCACGTCATACTAGCTGCTAGAGATGTCGTGATATTAGTGTGACGATGGTACGGACGAAGTACGAACGTGCATCAGAGAGGACCTGGAACCTGGTACCTGCTACCACCACGTCTCTGAC-

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Oligos for template synthesis:

- FPv2: CAGCTTGGACACCACGTCATAC GTCAGAGACGTGGTGGAGGAA
- CAGCTTGGACACCACGTCATACTAGCTGCTAGAGATGTGGTGATATTAGTGTGTGACGAT
- CAGCTTGGACACCACGTCATACGGAAGAAGACACAAAGACCTGATATTAGTGTGTGACGAT CAGCTTGGACACCACGTCATACTCAGGAGTCGAGAACTGAAGATATTAGTGTGTGACGAT
- CAGCTTGGACACCACGTCATACTGTGTACGTCAACACGTCAGATATTAGTGTGTGACGAT
- CAGCTTGGACACCACGTCATACCCATCCAACATCGTTGGAAGATATTAGTGTGTGACGAT

- CAGCTTGGACACCACGTCATACTCACGAAGCTGGATGATAGAGATATTAGTGTGTGACGAT CAGCTTGGACACCACGTCATACTAGCATCGATCGAACGTAGGATATTAGTGTGTGACGAT CAGCTTGGACACCACGTCATACTAGCATCGATCGAACGTAGGATATTAGTGTGTGACGAT
- GTCCTCTGATGCACGTTCGTACTTGTGCGTACCATCGTCACACACTAATATC
- GARCGTGCATCAGAGAGGACTCGACCACTGCAGGTGGAGCTCCAATTCCAGCTTCTAGGAAGACT GARCGTGCATCAGAGAGGACGTGCTTCCTCTGCTGCACCACGAATTCCAGCTTCTAGGAAGACT
- GAACGTGCATCAGAGAGGCCTGGTGTCGAGGTGAGCAGCAGCAATTCCAGCTTCTAGGAAGACT GAACGTGCATCAGAGAGGACTCGACGAGGTCCATCCTGGTCGCAATTCCAGCTTCTAGGAAGACT
- GAACGTGCATCAGAGAGGACCTGACACTGGTCGTGGTGGAGGCAATTCCAGCTTCTAGGAAGACT GAACGTGCATCAGAGAGGACCATCTCGACGACCTGCTCCTGGGAATTCCAGCTTCTAGGAAGACT GAACGTGCATCAGAGAGGACCACGAGGTCTCCACTGGTCCAGGAATTCCAGCTTCTAGGAAGACT
- -10:GAACGTGCATCAGAGAGGACCACTGAGCTGCTCCTCCAGGTGGAATTCCAGCTTCTAGGAAGACT
- GTCAGAGACGTGGTGGAGGAAGTCTTCCTAGAAGCTGGAATT

WO 2005/026387

PCT/DK2004/000630

Synthesis of identifier oligonucleotides:

ers FPv2 and RPv2 with TA= 53°C. The 160 bp products were gel-purified estimated from these measurements) were loaded on an agarose gel. fied on spectrophotometer. As a control, 20 ng of each of the templates (as using QIAquick Gel Extraction Kit from QIAGEN (Cat. No. 28706) and quantireactions each containing 0.05 pmol of the oligos Q-Temp1-X, Q-Temp2, Q-The 10 identifier oligonucleotides were assembled in 10 seperate 50 µl PCF Temp3-X and Q-Temp4 (x≃1 through 10) and 25 pmol of the external prim-

Preparation of samples for Q-PCR

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ies/5µl (0.00332 fmol/µl). $(160 \text{bp} \times 650 \text{ Da/bp} = 1.04 \times 105 \text{ g/mol}. 1 \text{ ng} = 9.615 \text{ fmol}). Diluted to <math>10^7 \text{ cop}$ prep. Volume was adjusted to 50 µl. Concentration: 4 ng/µl = 38.46 fmol/µl Sample A: Generated by mixing 20 ng from each identifier oligonucleotide

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5μl undil. Template #10 was mixed as follows:

Sample B: 20 ng/20µl stocks of each template were prepared. The sample

5µ1 2x dil. Template #9

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5μl 8x dil. Template #7 5µl 4x dit. Template #8

5µl 128x dil. Template #3 5µl 64x dil. Template #4 5µl 32x dil. Template #5

5µl 16x dil. Template #6

25

5μl 256x dil. Template #2

5µl 512x dil. Template #1

Concentration: $10 \text{ng}/50 \mu\text{l} = 0.20 \text{ ng}/\mu\text{l} = 1.923 \text{ fmol}/\mu\text{l}$. Diluted 579.2-fold to 107 copies/5µl (0.00332 fmol/µl).

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Sample A 116.55-fold to 10" copies/5 µl (0.33 fmol//µl) and subsequently per Standard curve: The samples for the standard curve was prepared by diluting

WO 2005/026387

PCT/DK2004/000630

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forming a 10-fold serial dilution of this sample. 5 µl was used for each PCR reaction. The standard curve is shown in Fig. 12.

Q-PCR reactions

- 5 For 5 ml premix (for one 96-well plate):
- 2.5 ml Taqman Universal PCR Master Mix (Applied Biosystems; includes Taq polymerase, dNTPs and optimized Taq pol. buffer)
- 450 µl RPv2 (10 pmol/ul)
- 25 µl Taqman probe (6-FAM-TCCAGCTTCTAGGAAGAC-MGBNFQ; 50 µM;
 - Applied Biosystems)

1075 µl H2O

40.5 µl premix was aliquoted into each well and 4.5 µl of relevant upstream PCR primer (FPv2 (for standard curve) or one of the codon specific primers

- 15 listed below; 10 pmol/µl) and 5 µl sample (H2O in wells for negative controls) was added. The codon-specific PCR primers were: (Tm calculations shown are from Vector NTI; matched to Tm for RPv2 (67.7°C))
- P1-1: GTCATACTAGCTGCAGAGATGTGGTGATA 66.8°C 20 P1-2: CATACGGAAGAGACAGAAGACGGAAGAGACAGAAGACTGATA 67.8°C
- P1-3: TCATACTCAGGAGTCGAGAACTGAAGATA 67.6°C P1-4: CATACTGTGTACGTCAACACGTCAGATA 67.4°C
- P1-5: CATACTGTGGAACTACCATCCAAGGATA 68.0°C
- P1-6: CCATCCAACATCGTTGGAAGAT 67.8°C 25 P1-7: CATACAACCTGTCCTGTGAGATCTGATA 67.7°C
- P1-7: CATACAACCIGIGAGAICIGAIA 67.7°C
 P1-8: ATACTCACGAAGCTGGATGATGAGATA 67.3°C
 P1-9: CATACTAGCATCGATCGAACGTAGGATA 68.1°C
 P1-10: TCATACTCGAAGCTACTGTCGAGATGATA 68.2°C
- P2-1: ATATTAGTGTGACGATGGTACGCA 67.8°C. 30 P3-1: ACAAGTACGAACGTGCATCAGAGA 67.7°C P4-1: CGAGCAGGACCTGGAACCT 67.7°C

68.3°C

P4-2: TCGACCACTGCAGGTGGA

	P4-3:	P4-3: GCTTCCTCTGCTGCACCA	66.7°C
	P4-4:	P44: GGTGTCGAGGTGAGCAGCA	69.1°C
	P4-5:	P4-5: CGACGAGGTCCATCCTGGT	08.6°C
	P4-6:	P4-6: GTGAGGAGCAGGTCCTCTGT	0.089
	P4-7:	P4-7: CTGACACTGGTCGTGGA	68.8°C
	P4-8:	P4-8: CATCTCGACGACCTGCTCCT	07.9°C
٠	P4-9:	P4-9: ACGAGGTCTCCACTGGTCCA	68.3°C
	P4-10:	P4-10: ACTGAGCTGCTCCTCCAGGT	66.5°C

PCT/DK2004/000630

5

WO 2005/026387

- 10 Thermocycling/measurement of fluoresence was performed on an Applied Biosystems ABI Prism 7900HT real-time instrument utilizing the standard cycling parameters:
- 95°C 10 min;
- 40 cycles of
- 95°C 15 sec;

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60°C 1 min

All samples were run in duplicate.

20 Result

Fig. 12 shows the standard curve calculated by the 7900HT system software. The log of the starting copy number was plotted against the measured C_T value. The relationship between C_T and starting copy number was linear in the range from 10 to 10⁸ template copies.

This standard curve was utilized by the system software to calculate the quantity in the "unknown" samples as shown below.

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PCT/DK2004/000630

PCT/DK2004/000630

13A				Expected	10000000	1000000	1000000	1000001	1000000	1000000	1000000	1000000	1000000	1000000	1000000	10000000	10000000	1000000	1000000	1000000	1000000	1000000	1000000	1000000	1000000	1000000	1000000
raphically in Fic				Observed B	11977503,00	480382,03	847478,56	948770,00	741304,40	1275155,50	1337928,50	747371,56	653874,00	705785,75	836037,90	14482606,00	12773780,00	1472576,80	2481824,80	2085476,40	1364621,40	2065813,60	1873777,20	1416153,00	1581067,00	1594593,80	1912277,40
Table I: Sample A (Shown graphically in Fig. 13A				Observed A	12539947,00	445841,90	884840,70	1013073,56	. 764187,94	1352874,60	1284075,60	658161,80	742187,20	824587,75	813550,75	13145159,00	13263911,00	1430704,80	2681652,00	1933106,80	1359684,40	2206709,80	1652718,10	1468208,10	1664467,50	1462520,60	2020088,20
Table I; Sa	Sample	. A: Equi-	molar	ratios	FPv2	P1-1	P1-2	P1-3	T 4	P1-5	9-1-6	P1-7	P1-8	P1-9	P1-10	P2-1	P3-1	P4-1	P4-2	P4-3	P4-4	P4-5	P4-6	P4-7	P4-8	P4-9	P4-10

Table II: Sample B (Shown graphically in Fig. 13B)

Observed Sample

Expected B: 2-fold Observed A

10000000 5,05E+06 10000000 10899,97 9765,625 19531,25 10000000 9765,625 19531,25 5000000 39062,5 374809,13 1250000 2500000 39062,5 312500 123734,13 156250 172005,64 625000 78125 53 44070,81 25419,85 166220,5 13469,12 1,72E+06 5,37E+06 5,09E+06 22733,17 39663,62 576151 70223,8 163687,44 646619,44 34748,89 156993,81 343176,78 1,49E+06 5,19E+06 (no signal) 110881,41 12732,32 4,97E+06 5,29E+06 42103,32 54480,62 .25542,8 9955,07 WO 2005/026387 P1-10 7-1-1 P1-2 P14 P1-5 P1-6 P1-9 P1-8 P2-1 P1-7 P4-1 P4-2 P4-3 P3-1

fold dynamic range, and reliable relative quantification of the tested codons in The results of the experiments show the possibility of accurately quantification of identifier oligonucleotides down to or even below 10 copies with a 9 various positions in the identifier oligonucleotide. . LO

691296,75 1250000

737661,44

316505,78

174134,64

156442,55 312500 283856,84 625000

115027,34 156250

137946,95

78125

43950,9

51293,07

2500000

,45E+06

1,42E+06

3,72E+06

While the invention has been described with references to specific methods changes may be made without departing from the invention. All patent and ilterature references cited herein are hereby incorporated by reference in and embodiments, it will be appreciated that various modifications and

their entirety. 2

The following Items describe embodiments of the first aspect of the present invention:

- tities, said encoded molecule being capable of forming part of a complex also Item 1: A method for obtaining structural information about an encoded molecule produced by a process comprising reaction of a plurality of chemical encomprising an identifier oligonucleotide containing codons informative of the identity of chemical entities which have participated in the formation of the encoded molecule, the method comprises 9 5
- a) mixing a primer oligonucleotide with the identifier oligonu-
- sion reaction to occur when the primer is sufficient complemenb) subjecting the mixture to a condition allowing for an extentary to a part of the identifier oligonucleotide, and

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- c) evaluating, based on measurement of the extension reaction, the presence, absence, or relative abundance of one or more
- Item 2: The method according to Item 1, wherein a composition of one, two, or more identifier oligonucleotides are processed simultaneously. 22

jecting a library of different complexes to a condition partitioning one or more Item 3: The method according to Item 2, the composition is a result of sub-

complexes having a certain property from the remainder of the library. ജ

WO 2005/026387

PCT/DK2004/000630

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Item 4: The method according to Item 1, wherein the condition which allows for an extension reaction to occur includes a polymerase or a ligase as well as suitable substrates.

Item 5: The method according to Item 4, wherein the condition includes a polymerase and a substrate comprising a blend of (deoxy)ribonucleotide riphosphates. ഗ

chemical entities are precursors for structural units appearing in the encoded Item 6: The method according to any of the preceding Items, wherein the molecule. 9

Item 7: The method according to any of the Items 1 to 6, wherein the process of producing the one or more encoded molecules comprises transferring one or more chemical entities to a nascent encoded molecule by a building block which further comprises an anti-codon.

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Item 8: The method of Item 7, wherein the information of the anti-codon is transferred in conjunction with the chemical entity to the nascent encoded molecule.

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Item 9: The method according to any of the preceding Items 1 to 8, wherein the identifier comprises two or more codons. Item 10: The method according to any of the preceding Items 1 to 8, wherein the identifier comprises three or more codons. 22

neighbouring codons of the identifier are spaced by a framing sequence. Item 11: The method according to any of the preceding Items, wherein

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WO 2005/026387

PCT/DK2004/000630

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Item 12: The method according to Item 11, wherein the framing sequence positions the reaction of a chemical entity in the synthesis history of the encoded molecule.

Item 13: The method according to any of the Items 1 to 12, wherein at least a part of the primer is complementary to a codon.

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Item 14: The method according to any of Items 1 to 13, wherein at least a part of the primer is complementary to a codon and an adjacent framing sequece.

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Item 15: The method according to any of the Items 1 to 13, wherein the codons have a length of four or more nucleotides.

15 Item 16: The method according to any of the Items 1 to 15, wherein the sequence comprising the codon and an adjacent framing sequence has a total length of 11 nucleotides or more.

Item 17: The method according to any of the Items 1 to 16, wherein the extension reaction is measured using the polymerase chain reaction (PCR), wherein the primer of Item 1 is involved in said PCR.

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Item 18: The method according to any of the Items 1 to 17, wherein a primer is labelled.

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Item 19: The method according to Item 18, wherein the primer is labelled with a small molecule, a radio active component, or a fluorogenic molecule.

Item 20: The method according to Item 19, wherein the small molecule label is selected from biotin, dinitrophenol, and digoxigenin, and the PCR amplicons are detected using an enzyme labelled streptavidin, anti-dinitrophenol. or anti-digoxigenin, respectively, reporter molecule.

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WO 2005/026387

PCT/DK2004/000630

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Item 21: The method according to any of the Items 1 to 19, wherein extension reaction is measured by real-time PCR.

Item 22: The method according to Item 21, wherein the real-time PCR involves the use of an oligonucleotide probe responsible for the generation of a detectable signal during the propagation of the PCR reaction.

Item 23: The method according to any of the Items 1 to 21, wherein the probe is designed to hybridise at a position downstream of a primer binding site.

Item 24: The method according to Item 22 or 23, wherein the probe is a 5' nuclease oligoprobe or a hairpin oligoprobe.

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Item 25. The method according to Items 2 or 3, wherein the Ilbrary comprises complexes with identifier oligonucleotides having n codon positions and the codons in said codon positions being selected from a set of m different codons.

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Item 26: The method according to Item 25, wherein a framing sequence is related to each of the n codon positions in a particular complex, sald framing sequence positions the reaction of a chemical entity in the synthesis history of the encoded molecule.

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Item 27: The method according to Item 25, wherein each codon in the set of m different codons differs from any other codons in the set in at least two nucleotide positions.

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Item 28: The method according to Item 26, wherein each framing sequence in a set of n different framing sequences differs from any other framing se-

30 quences in the set in at least two nucleotide positions.

Item 29: A method for identifying the chemical entities utilized in the formation of an encoded molecule or a composition of encoded molecules, wherein in separate compartments, n x m primers individually are mixed with an aliquot of a composition obtained by subjecting a library of different complexes to a condition partitioning said composition from the remainder of the library, subjected to a mixture of polymerase and substrate (deoxy)ribonucleotide triphosphates under conditions allowing for an extension reaction to occur when a primer is sufficient complementary to a part of one or more identifier oligonucleotides present in the aliquot, and evaluation, based on measurement of the extension reaction, the presence, absence, or relative abundance

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of one or more codons in each compartment.

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Item 30: A set comprising a collection of oligonucleotide primers, a polymerase, a composition of (deoxy)ribonucleotide triphosphates (dNTPs), and a library of complexes composed of a display molecule part and an identifier oligonucleotide, said oligonucleotide comprising codons informative of the identity of the chemical entities which has participated in the formation of the display molecule, wherein the oligonucleotide primers are sufficient complementary to codons appearing on the identifier oligo nucleotides in the library to allow for an extension reaction to occur.

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Item 31: An encoded molecule identified by a method according to Items 1 to 28.

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25 Method for Identifying a display molecule

The following pages will describe the second aspect of the invention: A METHOD FOR IDENTIFYING A DISPLAY MOLECULE, in which various patent and non-patent references cited in the present application are hereby incorporated by reference in their entirety. It is envisaged that any of the embodiments of the first aspect of the present invention may be used in combnation with any of the embodiments of the second aspect of the present invention, indeed any of the features described in relation to the first aspect of

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WO 2005/026387 PCT/DK2004/000630

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the present invention may be used in combination with any of the embodiments or features of the second aspect of the present invention, and vice versa.

- The second aspect of the present invention relates to a method for identifying from a library a display molecule having affinity towards one or more molecular targets. The display molecule is a part of a complex also comprising an identifier oligonucleotide that codes for said display molecule.
- Traditional drug discovery begins with a pathological phenomenon in an organism and the development of a therapeutic theory to combat this. A chemical concept follows to produce compounds for screening. Most of the processes for curing the pathological phenomenon originate with the understanding of some biological pathways and screening for an effect in tissues or cells. This may or may not eventually reveal a "target". The target can be identified by various conventional methods, including protein expressing, protein chemistry, structure-functional studies, knowledge of biochemical pathways, and genetic studies.
- In recent years, genetic information has increasingly guided the identification of molecular targets. These are derived from the knowledge of the genes of specific cell phenotypes that encode proteins that may be involved in the pathogenesis of a particular disease state.
- A lead is a compound, usually a small organic molecule that demonstrates a desired biological activity on a target. Usually, a collection of compounds, referred to as a "library", is screened before a useful lead is identified. Today, many libraries are commercially available or open to public. Most pharmaceutical companies house their own compilation of compounds that have been synthesised over several years and screened against a variety of tar-

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PCT/DK2004/000630

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been developing instrumentation capable of handling multiple micro titer plate ized plates and simplified assay formats all have made an impact on the relicompounds possible to screen simultaneously, different manufactures have ormats on the same platform using 384 and 1536-well plates. Advances in small volume liquid dispensing and pipetting, reliable handling of standardsands compounds simultaneously have been developed and are generally against the target. Techniques for handling the screening of several thoureferred to as high-throughput screening techniques. To push the limit of Each compound in a library must be screened by an appropriate assay ability of the high-throughput screening process.

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compounds has to be positioned in spatially discrete regions, usually in wells However, high-throughput screening has the disadvantage that each of the of a micro titer plate in order to observe an interaction with a target. If more pound displaying the appropriate biological activity. Thus, the full power of than a single compound is present, it is not feasible to discern which comcombinatorial chemistry cannot be applied because a collection of compounds usually is produced in a single container.

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oligonucleotide sequence is suitably a nucleic acid which identifies the poten-To be able to select a possible lead compound in a collection of compounds placed in the same container, libraries of bifunctional complexes have been evolved. Each bifunctional complex in the library comprises a potential lead compound coupled to an identifier oligonucleotide sequence. The identifier plexes can be eluated and the lead compound identified by sequencing the against a target, one or more of the potential leads may bind to the target. tial lead compound. When a library of bifunctional complexes is screened After removal of the remainder of the library, the binding bifunctional comidentifier oligonucleotide.

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Various techniques for producing bifunctional complexes are known from the prior art. Some attempts to form the complex comprising a molecule as well

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WO 2005/026387

PCT/DK2004/000630

as the identifier oligonucleotide that codes therefore, are based on the splitand-mix principle known from combinatorial chemistry, see e.g. WO 93/06121 A1, EP 643 778 B1, and WO 00/23458. Other attempts have focussed on the formation of encoded proteins using the The genetic information of the nucleic acid, usually mRNA or DNA, may not natural machinery of a cell and connecting the formed protein with the template nucleic acid that has coded for the amino acld components of the prolein. Examples of suitable systems are phage display, E. coli display, ribosome display (WO 93/03172), and protein-mRNA-fusions (WO 98/31700). Ŋ

for the formation of a new library enriched in respect of suitable binding prodentity of the chemical entities that has formed the protein because the nucleic acid can be amplified by known means, such as PCR, and processed necessarily be decoded between each round of selection to establish the 9 5

wherein the encoded molecule is not restricted to peptides and proteins. WO Recently, a method for encoding molecules has been suggested, which can be performed in several selection rounds without intermediate decoding,

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molecule connected to an identifier oligonucleotide coding for chemical entiprising a transferable chemical entity and an anticodon are initially provided. nealed together and the chemical entities are subsequently reacted to form segregated into a plurality of codons and a plurality of building blocks com-02/00419 and WO 02/103008 disclose methods for preparing virtually any ties which have reacted to form the display molecule. In short, a template Under hybridisation conditions, the template and building blocks are anthe motecute.

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display molecules having affinity towards a target using a library of bifunc-The present invention aims at providing an efficient method for identifying ional complexes.

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The second aspect of the present invention concerns a method for identifying display molecule(s) having affinity towards molecular target(s), comprising the steps of mixing one or more molecular target(s) associated with target oligonucleotide(s) and a library of bifunctional complexes, each bifunctional complex of the library comprising a display molecule attached to an identifier oligonucleotide, which codes for said display molecule,

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coupling to the target oligonucleotide(s) the identifier oligonucleotide of complexes comprising display molecules binding to the target, and

deducing the identity of the binding display molecule(s) and/or the molecular target(s) from the coupled product between the identifier oligonucleotide(s) and the target oligonucleotide(s).

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molecule and the coupling point on the identifier oligonucleotide sequence as selectivity, will increase with higher local concentration of ends of nucleotides gonucleotides, including appropriately selected lengths of target and identifier The second aspect of the present invention is based on the realization that a the target or the oligonucleotides. The proximity effect, and thus the power of close proximity of the identifier oligonucleotides relative to the target oligonucleotide compared to identifier oligonucleotides of complexes not comprising a display molecule having affinity toward the target. The tendency to be couwell as the length between the target molecule and the coupling point on the oligonucleotides, site of attachment of the target oligonucleotide to the target to be coupled together. Thus, various embodiments of the present invention may be envisaged to fine-tune the local concentration of the ends of the olitarget sequence, and (3) possible steric effects resulting from the nature of target and therefore, will be more prone to be coupled to the target oligonucleotide is obtained when a display molecule has binding affinity towards a pled together depends on various factors such as (1) the affinity of the displayed molecule towards the target, (2) the length between the displayed relative to the binding site of the display molecule, and size of target.

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PCT/DK2004/000630 WO 2005/026387

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effect, the target is usually expected to originate from an organism that harms bodiments the target may originate from a plant. In the quest for a compound certain disease. In the quest for discovering compounds with plant protective lected from human and animals, especially vertebras. However, in other em-The molecular target may be of a biological origin or may be a synthetic mowith therapeutical effect on the human or animal body, the target is usually expected to have an importance in a therapeutically theory that combats a lecular target. Typically, the molecular target stems from an organism sethe crop or a competing undesired plant. The organism may be a fungus

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The molecular target may be a protein, a small molecular hormone, a lipid, a polysaccharide, a whole cell, a nucleic acid, a metabolite, a heme group, etc. regulatory protein, a membrane channel or pump, a part of a signal transduction in the organism of being an enzyme, a hormone, a structural element, a in a preferred aspect the target is a protein. The protein may serve the funcdeleting one or more amino acids. 5

stemming from a biological origin may be derivatised by altering, adding, or

when a compound with fungicide effect is searched for or an insect when a

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compound having Insecticide effect is desired. Optionally, a protein target

two or more functionalities. Furthermore, the molecular target may be a solucomprise a prosthetic group. Also, the target may be a fusion protein having phosphatates, and proteases. The protein may occur as an independent enstituents occurring in the body or artificial components. In another preferred ble or insoluble agglomerate of one or more proteins and one or more subtity or may be dimers, trimers, tetramers, or polymers and the protein may ing cascade, an antibody, etc. Suitable target enzymes include kinases,

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embodiment, the molecular target is a nucleic acid, such as DNA or RNA aptamer or ribozyme. 23

The target may be immobilized to a support or be present as a solution or a emulsion, as appropriate. The target optionally immobilized on the support, may also form a stable or quasi-stable dispersion in the media. In a certain ဓ္က

PCT/DK2004/000630

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embodiment, the target is in solution and all the reactions occur in the solution too. The absence of an immobilization step reduces the background noise because there is no background surface to associate to. Thus the result of the assay may be more sensitive. In solution, the only background noise imaginable is when the oligonucleotides or display molecules of the library of complexes binds unspecific to the target molecules or the target oligonucleotides. The absence of an immobilization step generally necessitates a subsequent recovery step, eg. by chromatography.

In certain aspects of the invention, it is preferred to immobilize the molecular target on a solid support. The solid support may be beads of a column or the surface of a container. The immobilisation of the molecular target may ease the removal of the non-binding complexes by washing or similar means. In a certain embodiment, a cleavable linkage between the molecular target and the solid support is present. The cleavable linker is preferably selectively cleavable, that is, the linkage can be cleaved without cleaving other linkages in the target or the complexes. The cleavage of the linkage between the molecular target and the solid support may reduce the contribution from the background, such as complexes associated with the surface of the solid support and not binding to the molecular target.

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A single or a multitude of different molecular targets may be mixed with the library of complexes. If two or more different targets are mixed with the library of complexes it is appropriate to provide the molecular targets with a genetic sequence coding for the identity of the target in question. Proving the targets with identifying oligonucleotides allows for a simultaneous decoding of the binding partners, i.e. the molecular target and the display molecule. The simultaneous decoding is not only suitable for finding binding partners. It is also valuable for finding a possible cross-binding interaction or to find other display molecules competing for the same target. Furthermore, appropriate selection of display molecules and/or molecular targets, can generate useful

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WO 2005/026387

PCT/DK2004/000630

Information for preparing a structure-activity relationship (usually abbreviated

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The molecular target may be obtained in any suitable way. A variety of targets are commercially available, either as purified protein or as the corresponding cDNA. Other protein or peptide targets may be isolated from tissues or mRNA (or the corresponding cDNA) may be extracted from a tissue. Smaller peptides may be synthesised chemically using the standard solid—phase Fmoc peptide synthesis. When nucleic acids are used or included in the molecular target, it may be synthesised using the standard amedite synthesis method or by using the natural machinery.

The target can be associated with the target oligonucleotide using any suitable means and the association may involve a covalent or non-covalent linkagge. In an aspect of the invention the oligonucleotide is associated with the target utilizing a chemical synthesis. A protein usually comprises several groups that may be functionalized and used as attaching point. As examples the side chain of lysine contains an amino group, the side chain of serine contains a hydroxyl group and the side chain of cystein contains a thiol group, all of which may serve as anchoring point for a target oligonucleotide comprising e.g. a carboxylic acid group. In another aspect of the invention, the protein target is fused to a tag, such as a His-tag, Flag-tag, antibody, or streptavidin. The tag can be selectively recognized by an anti-body or small molecule such as biotin or dinitrophenol. The anti-body or the small molecule is attached to the target oligonucleotide, thereby ensumng an efficient coupling of the molecular target to the target oligonucleotide.

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The target oligonucleotide can be associated to the molecular target through a cleavable linker. The cleavable linkage can be used to separate the target oligonucleotide from the molecular target or the coupled product from the target at a point in time following the contacting between the target and Ilbrary of complexes. The target oligonucleotide can be distanced from the

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ble can be between the molecular target and the target oligonucleotide and at coding nucleotide sequence. A linker may remedy any interaction that possimolecular target by a suitable linker, such as a polyethylene linker or a nonthe same time provide suffice space for an enzyme to perform its action.

method is generally referred to as mRNA display. Optionally, the mRNA may A variety or methods for association of an oligonucleotide to a target is available for the skilled person in the art. An option involves the association of a target protein with the mRNA responsible for the formation thereof. This

be substituted with the corresponding cDNA. A method for generation such a single or a library of fusions between a protein and the mRNA responsible for their entirety by reference herein. The method of WO 98/31700 includes procodon operable linked to a protein encoding sequence, and a peptide accepstrand may be digested by RNase H. Another suitable method for generating tor at the 3' end and translating the protein encoding sequence to produce a the formation thereof is disclosed in WO 98/31700. The corresponding DNA a target Ilbrary is disclosed in WO 01/904.14, the content of which is incorpo-RNA-protein fusion. According to WO 00/32823 a DNA primer is covalently connected to the 3' end of the mRNA strand and extended by reverse tran-00/32823. The contents of both patent applications being incorporated in strand may be attached to the protein using the method disclosed in WO scriptase a to prepare the complementing DNA strand. The original RNA viding a RNA stand comprising a translation initiation sequence, a start rated herein by reference. 9 5

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get oligonucleotide is connected to the capsule via reactive groups positioned phage display, in which a target is displayed on the capsule of the phage and with a target using a method generally referred to as ribosome display. Ribosome display is disclosed in WO 93/03172, the content of which is included herein by reference. A still further option for association is a variation of the a target oligonucleotide is connected to the same capsule. Suitably, the tar-In accordance with another option, the target oligonucleotide is associated

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PCT/DK2004/000630 WO 2005/026387

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on proteins expressed on the capsule. A suitable reactive group is -SH emanating from cystein. A further option for associating the target with an oligonucleotide includes the sion of a target oligonucleotide connected to biotin and compartmentalization After the formation of the fusion protein in each compartment, the streptavidin method disclosed in M. Yonezawa et al, Nucleic acid research, 2003, vol. 31, part of the fusion protein binds to the biotin moiety of the target oligonucleo-No. 19 e118 (included by reference). The method includes the initial provithereof together with a transcription and translation system. The target oligonucleotide comprises a fusion gene coding for streptavidin and a target. tide, thereby associating the target with the target oligonucleotide. S 9

the function of a molecular target in the present invention and one or both the oligonucleotide sequence can be adhered to one or both the constant regions tamer or a library of aptamers comprising constant nucleic acid regions flankconstant region serves as target oligonucleotides. Alternatively, an additional In case the target is a nucleic acid, it may be of the aptamer type, i.e. an apthe same target but to different epitopes. The latter embodiment is of particuthe selection of pairs of aptamers that either binds to each other or binds to ing a random oligonucleotide part. The random oligonucleotide part serves to serve as the target oligonucleotide. The present invention also allows for lar relevance when evolving detection assays, where aptamers that bind to different epitopes on the same target may be desired.

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ated with the molecular target. The bifunctional complexes binding to the tarbe provided prior to or during with the mixing step. In case the target oligonubinding to the molecular target constitutes the target oligonucleotide associget and serving to associate an oligonuclectide to the molecular target may bifunctional complex may be a member of the library or may be a complex In certain embodiments, a bifunctional complex having a display molecule cleotide is associated with the molecular target during the mixing step the ဓ

erally saturated with the known display molecule prior to the mixing step. The procedures. To find a second or further binding compound, the target is genadded to the mixture. In some aspects of the invention, the display molecule known to bind to the target from the prior art or from preceding screening known binding molecule may or may not be attached to a nucleic acid seis a compound known to bind to the target. The display molecule may be

discovered simultaneously. The bifunctional complexes may bind to the same binding site of the molecular target or the bifunctional complexes may bind to a covalent or non-covalent chemical bond, e.g. by a technique known as click In case the target oligonucleotide emanates from an identifier oligonucleotide lional complexes are associated with a common molecular target and can be binding sites, the display molecules may be connected to each other through molecule they are usually adhered together after the identification process of in the library of complexes, two bifunctional complexes of a library of bifuncdifferent binding sites. In case the bifunctional complexes bind to the same nected through a disulphide bridge. In aspects of the invention in which the the present invention by a suitable linker in order to form the effective comchemistry. By way of example only, the two display molecules can be condisplay molecules binds to discrete binding domains on the same target

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chemical entities to form a second generation library, said second generation molecules with affinity towards a target is found using the present method or another method. In the event the affinity is in the lower range of what is desired, the initial display molecule is amended by reaction with one or more In a certain aspects of the invention an iterative method for finding the desired compound is applied. An initial display molecule or a pool of display library being used in the method of the present invention.

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WO 2005/026387

PCT/DK2004/000630

linker. By way of example only, blood factors, such as factor VIIa and Xa may separately subjected to the method of the invention, whereupon the Identified display molecules binding to the two or more targets are linked via a suitable compound having a part binding to a first blood factor and a part binding to a second blood factor. In other aspects of the invention, targets are linked by a in Nature, biochemical components interact. In a certain aspect of the prebe prevented or promoted in the their interaction with each other using a sent invention, two or more targets interacting in a biological context are

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suitable linker, as disclosed in Fig. 10.

A library of molecular targets may be generated by starting out from a library of DNA molecules, usually cDNA molecules, and preparing the correspondaspect of the invention, the mixture step therefore includes that a molecular ing RNA strands by a suitable RNA polymerase. In according with a certain target library comprising different peptides each attached to the nucleic responsible for the formation thereof is mixed with a library of complexes.

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The coupling of the target oligonucleotide and the identifier oligonucleotide is are relatively remote from the target oligonucleotide and the tendency of the duced. Among other things, the unspecific coupling depends on the concenpromoted due to the relative high local concentration of the ends of oligonucleotides. The complexes in solution, i.e. complexes not bund to the target, target oligonucleotides to be coupled to such unspecific complexes is reration of the complexes in solution.

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ter. In case the target Is in solution or made in solution by cleavage of a bond performed by adding a suitable buffer and the removal may be preformed by ained on a filter while non-binding complexes is transported through the fil-In an aspect of the invention the non-binding library members are either diuted or at least partly removed prior to the coupling step. Dilution may be event the target molecule is immobilized on a bead, the beads may be rewashing one or more times with a suitable liquid, such as a buffer. In the ဓ

moval can be performed by chromatography, such as size-exclusion chromaimmobilising the target, optional following one or more washing step, the retography.

moval of non-binding library members prior to the coupling of the target oli-Thus, in a certain aspect of the invention, the mixing step includes the regonucleotide and the identifier nucleotide together.

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tain nucleic acids may perform a binding interaction or a catalytical action on double stranded form during the contacting with the molecular target, as certhe components present during the mixing step. Thus, in one embodiment of It may be an advantage to have all or at least a part of the nucleotides on a the invention, the target oligonucleotide and/or the identifier oligonucleotide partly or fully is hybridised to a complementing oligonucleotide.

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lected from the group consisting of chemical means, enzymatic means, and The coupling may be performed using any suitable means that ensures a physical connection. Suitably, the coupling is performed using means sedesign means.

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The chemical means for coupling the ends together can be selected from a large plethora. Suitable examples include that

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gonucleotide end comprises a 5'-phosphor-2-imidazole group. When reacted a) a first oligonucleotide end comprises a 3'-OH group and the second olia phosphodiester internucleoside linkage is formed,

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b) a first oligonucleotide end comprising a phosphoimidazolide group and the 3'-end and a phosphoimidazolide group at the 5'-and. When reacted together a phosphodiester internucleoside linkage is formed,

second oligonucleotide comprising a 5'-iodine. When the two groups are rec) a first oligonucleotide end comprising a 3'-phosphorothloate group and a acted a 3'-O-P(=O)(OH)-S-5' internucleoside linkage is formed, and ဓ

WO 2005/026387

PCT/DK2004/000630

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d) a first oligonucleotide end comprising a 3'-phosphorothioate group and a second oligonucleotide comprising a 5'-tosylate. When reacted a 3'-O-P(=0)(OH)-S-5' internucleoside linkage is formed.

- Suitably, the target oligonucleotide or a complementing target oligonucleotide chemical reaction strategy for the coupling step generally includes the formation of a phosphodiester intemucleoside linkage. In accordance with this asand the indentifier oligonucleotide or a complementing identifier oligonucleoide operatively are joined together, so that as to allow a nucleic acid active embodiment, the coupling is performed so as to allow a polymerase to recenzyme to recognize the coupling area a substrate. Notably, in a preferred ognise the coupled strand as a template. Thus, in a preferred aspect, a pect, method a) and b) above is preferred. S 6
- enzymatic means is in general selected from the enzymes of the polymerase reaction is specific, i.e. the risk of side reactions in virtually not present. The The enzymatic means is in some instances preferred because the coupling type, ligase type, and restriction enzymes, as well as any combination

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Faq DNA ligase, T4 DNA ligase, T7 DNA ligase, and E. coli DNA ligase. The choice of the ligase depends to a certain degree on the design of the ends to ferred, while a Taq DNA ligase may be preferred for a sticky end ligation, i.e. a ligation in which an overhang on each end is a complement to each other. Ligases are useful means for the coupling step. Suitable examples include be joined together. Thus, if the ends are blunt, T4 DNA ligase may be pre-

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abut each other a ligase can ligate the ends together. If a gap exists between In a certain aspect of the invention, a connector oligonucleotide is used. The target oligonucleotide and a region complementing a distal part of the identifier oligonucleotide. If the ends of the target and identifier oligonucleotides connector oligonucleotide has a region complementing a distal part of the

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the ends, a polymerase may be used to fill the gap and a ligase may subsequently perform a ligation. The regions of the connector oligonucleotide complementing the identifier oligonucleotide and target oligonucleotide, respectively, may independently be chose, e.g. in the range of 6 to 16 nucleotides, preferably in the range of 8 to 12 nucleotides. In a particular aspect of the invention the connector oligonucleotide is added in excess relative to the total target and identifier oligonucleotides to saturate the ends of complexes not bound to a target.

In another aspect of the invention the coupling is performed by design means. As an example, the regions at the distal ends of the target and identifier oligonucleotides are designed to be complementary to each other. Under hybridisation conditions polymerase is then allowed to extend the target oligonucleotide as well as the identifier oligonucleotide to obtain a double stranded coupled product.

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Still another coupling means include the design of the ends of one or more oligonucleotides with sticky ends. In a certain aspect, the target oligonucleotide and/or the identifier oligonucleotide is provided with a sticky end to allow a ligase or a polymerase or a mixture thereof to adjoin the oligonucleotides. Suitably, the sticky ends can be formed by a restriction nuclease. In a practical approach, the target and the identifier oligonucleotides are initially double stranded and provide at the ends with a restriction site. Following the initial contact, the mixture is treated with a restriction nuclease to form the sticky ends. After subsequent removal of the restriction enzyme from the mixture, a ligase is allowed to perform its enzymatic action to form a ligation product.

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The coupled product of the identifier and target oligonucleotides comprises the information necessary for decoding the identity of the display molecule and optionally also of the molecular target. The coupled product may be analysed directly in some instances to reveal the identity of the display molecules that have performed an interaction with the molecular target. As an example,

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WO 2005/026387

PCT/DK2004/000630

the coupled product may be detached from the target-display molecule interaction using a cut by a restriction nuclease at positions of the coupled product flanking the informative part. The informative part can be decoded in a standard sequencing machine. In general however, it is preferred to include the informative part of the coupled product in to a suitable vector and transfer the vector to a host organism. The host organisms may then be cultivated on a suitable substrate and allowed to form colonies. Samples from the colonies may be used for sequencing in a sequencing machine.

cal to the coupled product. The extension product or the amplification product priately, the extended strand is designed with another priming site which may may be analysed directly or be incorporated into a vector which subsequently lymerase to extent the primer using the coupled product as template. Approquences complementary thereto at the proximal end Is provided with a primmerase to perform an extension of the primer to produce a sequence Identiof the invention, the coupled oligonucleotide is amplified by PCR using primis transformed into a host organism as explained above. In a certain aspect ing site. The priming site may be used for annealing a primer to allow a poin another approach, the target and/or the identifier oligonucleotides or seallow a second (reverse) primer to anneal thereto and subsequently a polyng sites positioned proximal to the display molecule and the molecular target, respectively, and flanking the informative part of the coupled product. **`**2 5 8

Following the coupling step, the target-display molecule conjugate may be recovered. Any method that result in a recovery may in principle be used, including filtering, washing, elution, chromatography, etc. In a preferred aspect the target-display molecule conjugate Is recovered by chromatography following the coupling of the target and the Identifier oligonucleotides. Optionally, the various recovering methods may be combined. As an example, a washing or elution step may precede the chromatography step. A presently preferred chromatography method is size-exclusion chromatography. The chromatography is usually performed on a sample in which the target-display

molecule conjugate is in solution. In one aspect of the invention, the target has been cleaved from a solid support prior to the chromatography step.

oligonucleotides prior to amplification. Usually, the cleavage is preceded of a In an aspect of the invention, selective cleavable chemical moieties at each step removing the non-binding complexes. The liberated product may be reend of the coupled oligonucleotides are cleaved to liberate the coupled covered as described above and subjected to amplification.

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linkage. In an embodiment of the invention, the target oligonucleotide and the identifier oligonucleotide are coupled together, whereby a product is obtained wherein a display molecule is bound to the molecular target and the identifier said molecule. Usually, the display molecule part of the complex is bound to The invention also pertains to a conjugate comprising a molecular target asthe target. The target oligonucleotide and/or the Identifier oligonucleotide of coupled oligonucleotide is amplifiable. The amplifiability is usually obtained display molecule attached to an identifier oligonucleotide, which codes for sociated with an oligonucleotide and a bifunctional complex comprising a and/or the display molecule, respectively, through a selectively cleavable and target oligonucleotides are coupled together. In a certain aspect, the the conjugate are in a certain embodiment joined to the molecular target through a coupling that involves a phospodiester internucleoside linkage.

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The present invention also extends to a display molecule identified by any of the methods disclosed herein. 23

Detailed Description

Complex

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The complex used in the present invention comprises a display molecule and an identifier oligonucleotide. The identifier oligonucleotide comprises identifygonucleotide identifies the molecule uniquely, I.e. in a library of complexes a ing moieties that identifies the display molecule. Preferably, the identifier oli-

WO 2005/026387

75

PCT/DK2004/000630

particular identifier oligonucleotide is capable of distinguishing the molecule it is attached to from the rest of the display molecules.

rectly to each other or through a bridging moiety. In one aspect of the inven-The display molecule and the identifier oligonucleotide may be attached diion, the bridging molety is a selectively cleavable linkage. Ŋ

which is amplifiable, Identifier oligonucleotides comprising a sequence of nu-The method may in certain embodiments be performed without amplification amount of separated coupled product oligonucleotide is relatively low, it is in general preferred to use an identifier oligonucleotide and a coupled prodct after the coupling step. However, when larger libraries are used and the cleotides may be amplified using standard techniques, like PCR.

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of the molecule. When the identifier oligonucleotide comprises more than one quence of codons can be decoded to identify reactants used in the formation codon, each member of a pool of chemical entities can be identified and the order of codons is informative of the synthesis step each member has been The identifier oligonucleotide may comprise two or more codons. The seincorporated in.

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length. The codon may be a single nucleotide or a plurality of nucleotides. In some aspects of the invention, it is preferred that each codon independently comprises four or more nucleotides, more preferred 4 to 30 nucleotides. The sequence of the nucleotides in each codon may have any sultable

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suitable framing sequence. Preferably, all or at least a majority of the codons be separated by a framing sequence. Depending on the encoded molecule formed, the identifier oligonucleotide may comprise further codons, such as 3, 4, 5, or more codons. Each of the further codons may be separated by a ranged in sequence, I.e. next to each other. Two neighbouring codons may The identifier oligonucleotide will in general have at least two codons ar-

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of the identifier oligonucleotide are separated from a neighbouring codon by a framing sequence. The framing sequence may have any suitable number of nucleotides, e.g. 1 to 20. Alternatively, codons on the identifier oligonucleotide may be designed with overlapping sequences.

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The framing sequence, if present, may serve various purposes. In one setup plate with an anti-codon will occur in frame. Moreover, the framing sequence affinity. The high affinity region may ensure that the hybridisation of the temcodons. In another setup, the frames have alternating sequences, allowing of the invention, the framing sequence identifies the position of the codon. for addition of building blocks from two pools in the formation of the library. Usually, the framing sequence either upstream or downstream of a codon The framing sequence may also or in addition provide for a region of high comprises information which allows determination of the position of the may adjust the annealing temperature to a desired level. A framing sequence with high affinity can be provided by incorporation of one affinity, such as 2'-0-methyl substitution of the ribose moiety, peptide nucleic sine. Alternatively, or in addition, the framing sequence may be subjected to backbone modification. Several back bone modifications provides for higher acids (PNA), and 2'-4' O-methylene cyclisation of the ribose moiety, also rebase. Examples of nucleobases having this property are guanine and cytoor more nucleobases forming three hydrogen bonds to a cognate nucleoferred to as LNA (Locked Nucleic Acid).

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The identifier oligonucleotide may comprise one or two flanking regions. The plex or the flanking region may comprise a label that may be detected, such flanking region can encompass a signal group, such as a flourophor or a radio active group to allow for detection of the presence or absence of a comas biotin. When the identifier oligonucleotide comprises a biotin moiety, the dentifier oligonucleotide may easily be recovered.

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WO 2005/026387

PCT/DK2004/000630

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ments comprise an affinity region having the property of being able to hybrid-The flanking region(s) can also serve as priming sites for amplification reactions, such as PCR. The identifier oligonucleotide may in certain embodiise to a building block.

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the sense or the anti-sense format, i.e. the identifier oligonucleotide can be a It is to be understood that when the term identifier oligonucleotide is used in quence complementary thereto. Moreover, the identifier oligonucleotide may the present description and claims, the identifier oligonucleotide may be in sequence of codons which actually codes for the molecule or can be a sebe single-stranded or double-stranded, as appropriate.

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of having an effect on the target. When the target is of pharmaceutical importance, the molecule is generally a possible drug candidate. The complex may other embodiment of the invention, the molecule is encoded, I.e. formed by a molecule. Optionally, this reaction product may be post-modified to obtain the The display molecule part of the complex is generally of a structure expected the cleavage of one or more chemical bonds attaching the encoded molecule to the indentifier in order more efficiently to display the encoded molecule. In final molecule displayed on the complex. The post-modification may involve the natural machinery, such as the methods disclosed in WO 92/02536, WO still another embodiment the display molecule is a polypeptide formed using tag, e.g. a nucleic acid tag identifying each possible drug candidate. In anbe formed by tagging a library of different possible drug candidates with a variety of reactants which have reacted with each other and/or a scaffold

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91/05058, and US 6,194,550.

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The formation of a synthetic encoded molecule generally starts by a scaffold, i.e. a chemical unit having one or more reactive groups capable of forming a connection to another reactive group positioned on a chemical entity, thereby generating an addition to the original scaffold. A second chemical entity may react with a reactive group also appearing on the original scaffold or a

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encoded molecule may be mediated by a bridging molecule. As an example, amine group a connection between these can be mediated by a dicarboxylic display molecule is not produced using the naturally translation system in an naturally occurring or an artificial substance. In an aspect of the invention, a if the nascent encoded molecule and the chemical entity both comprise an in vitro process. In other aspects of the invention, the display molecule is a entities may be involved in the formation of the final reaction product. The reactive group incorporated by the first chemical entity. Further chemical formation of a connection between the chemical entity and the nascent acid. A display molecule is in general produced in vitro and may be a polypeptide produced using the natural translation machinery.

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generally comprises an anti-codon. In some embodiments the building blocks prior to the participation in the formation of the reaction product leading to the eliminations of the encoded molecule may be attached to a building block also comprise an affinity region providing for affinity towards the nascent final display molecule. Besides the chemical entity, the building block The chemical entities that are precursors for structural additions or complex

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molecule by a building block, which further comprises an anticodon. The antihowever, it is important that a correspondence is maintained in the complex. The chemical entities are preferably reacted without enzymatic interaction in some aspects of the invention. Notably, the reaction of the chemical entities into a protein using a tRNA loaded with a natural or unnatural amino acid. In is preferably not mediated by ribosomes or enzymes having similar activity. In another aspect of the invention a ribosome is used to translate an mRNA transfer of genetic information and chemical entity may occur in any order, Thus, the chemical entities are sultably mediated to the nascent encoded still another aspect of the invention, enzymes having catalytic activities codon serves the function of transferring the genetic information of the building block in conjunction with the transfer of a chemical entity. The

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PCT/DK2004/000630 WO 2005/026387

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different from that of ribosomes are used in the formation of the display molecule

genetic information of the anti-codon to the nascent complex by an extension acid template. Another method for transferring the genetic information of the According to certain aspects of the invention the genetic information of the anti-codon is transferred by specific hybridisation to a codon on a nucleic complex, e.g. by ligation. A still further method involves transferring the complementary to the anti-codon and attach this oligonucleotide to the anti-codon to the nascent complex is to anneal an oligonucleotide eaction using a polymerase and a mixture of dNTPs.

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The chemical entity of the building block may in certain cases be regarded as a nascent encoded molecule it is to be understood that not necessarily all the pears on the nascent encoded molecule. Especially, the cleavage resulting in present application with claims is stated that a chemical entity is reacted with a precursor for the structural entity eventually incorporated into the encoded molecule. In other cases the chemical entity provides for the ellminations of encoded molecule. Also, as a consequence of the reactions involved in the atoms of the original chemical entity is to be found in the eventually formed connection, the structure of the chemical entity can be changed when it apchemical units of the nascent encoded molecule. Therefore, when it in the quent step can participate in the formation of a connection between a nasthe release of the entity may generate a reactive group which in a subsecent complex and a chemical entity.

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ween the chemical entity of the building block and another chemical entity or groups which appears on the chemical entity is suitably one to ten. A building group capable of participating in a reaction which results in a connection beblock featuring only one reactive group is used i.a. in the end positions of The chemical entity of the building block comprises at least one reactive a scaffold associated with the nascent complex. The number of reactive

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polymers or scaffolds, whereas building blocks having two reactive groups are suitable for the formation of the body part of a polymer or scaffolds capable of being reacted further. One, two or more reactive groups intended for the formation of connections, are typically present on scaffolds. Non-limiting examples of scaffolds are opiates, steroids, benzodiazepines, hydantoines, and peptidylphosphonates.

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The reactive group of the chemical entity may be capable of forming a direct connection to a reactive group of the nascent complex or the reactive group of the building block may be capable of forming a connection to a reactive group of the nascent complex through a bridging fill-in group. It is to be understood that not all the atoms of a reactive group are necessarily maintained in the connection formed. Rather, the reactive groups are to be regarded as precursors for the structure of the connection.

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The subsequent cleavage step to release the chemical entity from the building block can be performed in any appropriate way. In an aspect of the invention the cleavage involves usage of a chemical reagent or an enzyme. The cleavage results in a transfer of the chemical entity to the nascent encoded molecule or in a transfer of the nascent encoded molecule to the chemical entity of the building block. In some cases it may be advantageous to introduce new chemical groups as a consequence of linker cleavage. The new chemical groups may be used for further reaction in a subsequent cycle, either directly or after having been activated. In other cases it is desirable that no trace of the linker remains after the cleavage.

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In another aspect, the connection and the cleavage is conducted as a simultaneous reaction, i.e. either the chemical entity of the building block or the nascent encoded molecule is a leaving group of the reaction. In some aspects of the invention, it is appropriate to design the system such that the connection and the cleavage occur simultaneously because this will reduce the number of steps and the complexity. The simultaneous connection and

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WO 2005/026387

81

PCT/DK2004/000630

cleavage can also be designed such that either no trace of the linker remains or such that a new chemical group for further reaction is introduced, as described above.

The attachment of the chemical entity to the building block, optionally via a suitable spacer can be at any entity available for attachment, e.g. the chemical entity can be attached to a nucleobase or the backbone. In general, it is preferred to attach the chemical entity at the phosphor of the internucleoside linkage or at the nucleobase. When the nucleobase is used for attachment of the chemical entity, the attachment point is usually at the 7 position of the purines or 7-deaza-purins or at the 5 position of pyrimidines. The nucleotide may be distanced from the reactive group of the chemical entity by a spacer moiety. The spacer may be designed such that the conformational space sampled by the reactive group is optimized for a reaction with the reactive

group of the nascent encoded molecule.

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The display molecules of the invention may have any chemical structure. In a preferred aspect, the display molecule can be any compound that may be synthesized in a component-by-component fashion. In some aspects the display molecule is a linear or branched polymer. In another aspect the display molecule is a scaffolded molecule. The term "display molecule" also comprises naturally occurring molecules like a-polypeptides etc, however produced *in vitro* usually in the absence of enzymes, like ribosomes. In certain aspects, the display molecule of the library is a non-a-polypeptide.

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The display molecule may have any molecular weight. However, in order to be orally available, it is in this case preferred that the display molecule has a molecular weight less than 2000 Daltons, preferably less than 1000 Dalton, and more preferred less than 500 Daltons.

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The size of the library may vary considerably pending on the expected result of the inventive method. In some aspects, it may be sufficient that the library

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some aspects, the library comprises 1,000 or more different complexes, more more than two different complexes are desired to obtain a higher diversity. In preferred 1,000,000 or more different complexes. The upper limit for the size of the library is only restricted by the size of the vessel in which the library is comprised. It may be calculated that a vial may comprise up to 1014 different comprises two, three, or four different complexes. However, in most events, complexes.

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Methods for forming libraries of complexes

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herein by reference as well as methods of the present applicant not yet public available, including the methods disclosed in DK PA 2002 01955 filed 19 Demolecule part of the complex may be formed by a variety of processes. Genmethods may be used, and the entire content of the patent applications are The complexes comprising an identifier oligonucleotide having two or more erally, the preferred methods can be used for the formation of virtually any kind of encode molecule. Suitable examples of processes include prior art codons that codes for reactants that have reacted in the formation of the 02/074929, and WO 02/103008, the content of which being incorporated cember 2002, and DK PA 2003 00430 filed 20 March 2003. Any of these methods disclosed in WO 93/20242, WO 93/06121, WO 00/23458, WO included herein by reference.

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polymerase to incorporate unnatural nucleotides as building blocks. Initially, a derivatives, the chemical entities are reacted to form a reaction product. The annealed to each of the templates and a polymerase is extending the primer plurality of template oligonucleotides is provided. Subsequently primers are Subsequent to or simultaneously with the incorporation of the nucleotide encoded molecule may be post-modified by cleaving some of the linking Below four preferred embodiments are described. A first embodiment using nucleotide derivatives which have appended chemical entities. disclosed in more detail in WO 02/103008 is based on the use of a moieties to better present the encoded molecule.

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WO 2005/026387

PCT/DK2004/000630

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Several possible reaction approaches for the chemical entities are apparent.

First, the nucleotide derivatives can be incorporated and the chemical entities subsequently polymerised. In the event the chemical entities each carry two or bridging molety. Exemplary of this approach is the linking of two chemical nucleotide building block, such as an ester or a thioester group. An adjacent interspaced reactive group to obtain a linkage to the chemical entity, e.g. by reactive groups, the chemical entities can be attached to adjacent chemical bond. Adjacent chemical entities can also be linked together using a linking groups are amine and carboxylic acid, which upon reaction form an amide approach is the use of a reactive group between a chemical entity and the entities each bearing an amine group by a bi-carboxylic acid. Yet another entities by a reaction of these reactive groups. Exemplary of the reactive building block having a reactive group such as an amine may cleave the an amide linking group.

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hybridisation of building blocks to a template and reaction of chemical entities blocks, wherein each building block comprises an anti-codon and a chemical i.e. a codon, on the template. Subsequent to the annealing of the anti-codon approach comprises that templates are contacted with a plurality of building entity. The anti-codons are designed such that they recognise a sequence, A second embodiment for obtainment of complexes pertains to the use of attached to the building blocks in order to obtain a reaction product. This and the codon to each other a reaction of the chemical entity is effected.

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reaction of the reactive group of the chemical entity may be effected at any The template may be associated with a scaffold. Building blocks bringing chemical entities in may be added sequentially or simultaneously and a lime after the annealing of the building blocks to the template.

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enzymatical ligation of building blocks when these are lined up on a template. A third embodiment for the generation of a complex includes chemical or

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templates are contacted with building blocks comprising anti-codons linked to subsequently ligated to each other and a reaction of the chemical entities is effected to obtain a reaction product. The method is disclosed in more detail chemical entities. The two or more anti-codons annealed on a template are initially, templates are provided, each having one or more codons. The in DK PA 2003 00430 filed 20 March 2003.

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complex comprising a scaffold and an affinity region is annealed to a building Subsequently the anti-codon region of the building block is transferred to the the anti-codon. This method is disclosed in detail in DK PA 2002 01955 filed be transferred prior to, simultaneously with or subsequent to the transfer of nascent complex by a polymerase. The transfer of the chemical entity may building block to the nascent complex. The method implies that a nascent A fourth embodiment makes use of the extension by a polymerase of an affinity sequence of the nascent complex to transfer the anti-codon of a block comprising a region complementary to the affinity section. 19 December 2002 and DK PA 2003 01064, filed 11 July 2003.

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Thus, the codons are either pre-made into one or more templates before the encoded molecules are generated or the codons are transferred simultaneously with the formation of the encoded molecules.

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After or simultaneously with the formation of the reaction product some of the linkers to the template may be cleaved, however at least one linker must be maintained to provide for the complex

Nucleotides

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sequence of nucleotides, i.e. an oligonucleotide. Each nucleotide monomer is bone. The back bone may in some cases be subdivided into a sugar moiety normally composed of two parts, namely a nucleobase moiety, and a back-The nucleotides used in the present invention may be linked together in a and an internucleoside linker.

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WO 2005/026387

85

PCT/DK2004/000630

hydroxy-5-methyl-4-triazolopyridine, isocytosine, isoguanine, inosine and the cleobases are adenine, guanine, thymine, cytosine, uracil, purine, xanthine, diaminopurine, 8-oxo-N⁶-methyladenine, 7-deazaxanthine, 7-deazaguanine, N⁴,N⁴-ethanocytosin, N⁶,N⁸-ethano-2,6-diamino-purine, 5-methylcytosine, 5- $(\mathrm{C}^3\mathrm{-C}^6)$ -alkynylcytosine, 5-fluorouracil, 5-bromouracil, pseudoisocytosine, 2-The nucleobase moiety may be selected among naturally occurring nucleobases as well as non-naturally occurring nucleobases. Thus, "nucleobase" No. 5,432,272. The term "nucleobase" is intended to cover these examples includes not only the known purine and pyrimidine hetero-cycles, but also heterocyclic analogues and tautomers thereof. Illustrative examples of nu-"non-naturally occurring" nucleobases described in Benner et al., U.S. Pat Ω 9

bases are adenine, guanine, thymlne, cytosine, 5-methyicytosine, and uracil, which are considered as the naturally occurring nucleobases in relation to therapeutic and diagnostic application in humans.

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as well as analogues and tautomers thereof. Especially interesting nucleo-

Examples of suitable specific pairs of nucleobases are shown on p27 of this application, entitled "Natural base pairs", "synthetic base pairs", "synthetic purine bases pairing with natural pyrimidines".

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this application (B denotes a nucleobase). The sugar molety of the backbone Suitable examples of backbone units are shown diagrammatically on p28 of ribose (LNA). Suttably the nucleobase is attached to the 1' position of the is sultably a pentose but may be the appropriate part of an PNA or a sixmember ring. Suitable examples of possible pentoses include ribose, 2.deoxyribose, 2'-0-methyl-ribose, 2'-flour-ribose, and 2'-4'-0-methylenepentose entity.

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An internucleoside linker connects the 3' end of preceding monomer to a 5' end of a succeeding monomer when the sugar molety of the backbone is a pentose, like ribose or 2-deoxyribose. The internucleoside linkage may be

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internucleoside linker can be any of a number of non-phosphorous-containing phosphoramidate, phosphotriester, and phosphodithioate. Furthermore, the the natural occurring phospodiester linkage or a derivative thereof. Examples of such derivatives include phosphorothioate, methylphosphonate, linkers known in the art.

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and inosine. Inosine is a non-specific pairing nucleoside and may be used as compounds depicted on p30 of this application, entitled "Examples of Univermembers of the RNA family include adenosine, guanosine, uridine, cytidine, forming part of the DNA as well as the RNA family connected through phosoxyadenosine, deoxyguanosine, deoxythymidine, and deoxycytidine. The universal base because inosine can pair nearly isoenergetically with A, T, and C. Other compounds having the same ability of non-specifically base-Preferred nucleic acid monomers include naturally occurring nucleosides pairing with natural nucleobases have been formed. Suitable compounds which may be utilized in the present invention includes among others the phodiester linkages. The members of the DNA family include desal Bases",

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Building block 2

the participation in the formation of the reaction product leading the final en-The chemical entities that are precursors for structural additions or eliminalions of the encoded molecule may be attached to a building block prior to coded molecule. Besides the chemical entity, the building block generally comprises an anti-codon.

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tween the chemical entity of the building block and another chemical entity or a scaffold associated with the nascent complex. The connection is facilitated group capable of participating in a reaction which results in a connection beby one or more reactive groups of the chemical entity. The number of reac-The chemical entity of the building block comprises at least one reactive live groups which appear on the chemical entity is suitably one to ten. A

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WO 2005/026387

PCT/DK2004/000630

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folds capable of being reacted further. One, two or more reactive groups inbuilding block featuring only one reactive group is used i.a. in the end positions of polymers or scaffolds, whereas building blocks having two reactive groups are suitable for the formation of the body part of a polymer or scafended for the formation of connections, are typically present on scaffolds.

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derstood that not all the atoms of a reactive group are necessarily maintained connection to a reactive group of the nascent complex or the reactive group in the connection formed. Rather, the reactive groups are to be regarded as The reactive group of the building block may be capable of forming a direct group of the nascent complex through a bridging fill-in group. It is to be unof the building block may be capable of forming a connection to a reactive precursors for the structure of the connection.

ing block can be performed in any appropriate way. In an aspect of the invenor in a transfer of the nascent encoded molecule to the chemical entity of the groups may be used for further reaction in a subsequent cycle, either directly The subsequent cleavage step to release the chemical entity from the buildresults in a transfer of the chemical entity to the nascent encoded molecule or after having been activated. In other cases it is desirable that no trace of tion the cleavage involves usage of a reagent or an enzyme. The cleavage chemical groups as a consequence of linker cleavage. The new chemical building block. In some cases it may be advantageous to introduce new the linker remains after the cleavage.

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complexity. The simultaneous connection and cleavage can also be designed nascent encoded molecule is a leaving group of the reaction. In general, it is In another aspect, the connection and the cleavage is conducted as a simuloccur simultaneously because this will reduce the number of steps and the taneous reaction, i.e. either the chemical entity of the building block or the preferred to design the system such that the connection and the cleavage

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such that either no trace of the linker remains or such that a new chemical group for further reaction is introduced, as described above.

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linkage or at the nucleobase. When the nucleobase is used for attachment of preferred to attach the chemical entity at the phosphor of the internucleoside sultable spacer can be at any entity available for attachment, e.g. the chemipurines or 7-deaza-purins or at the 5 position of pyrimidines. The nucleotide cal entity can be attached to a nucleobase or the backbone. In general, it is may be distanced from the reactive group of the chemical entity by a spacer The attachment of the chemical entity to the building block, optionally via a the chemical entity, the attachment point is usually at the 7 position of the sampled by the reactive group is optimized for a reaction with the reactive molety. The spacer may be designed such that the conformational space group of the nascent encoded molecule or reactive site.

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codon. The anticodon may be adjoined with a fixed sequence, such as a se-The anticodon complements the codon of the identifier oligonucleotide sequence and generally comprises the same number of nucleotides as the quence complementing a framing sequence.

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Various specific building blocks are envisaged. Building blocks of particular interest are shown below

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Building blocks transferring a chemical entity to a recipient nucleophilic group The building block indicated below is capable of transferring a chemical entity ring serves as an activator, i.e. a labile bond is formed between the oxygen lower horizontal line illustrates the building block and the vertical line illustrates a spacer. The 5-membered substituted N-hydroxysuccinimid (NHS) (CE) to a recipient nucleophilic group, typically an amine group. The bold atom connected to the NHS ring and the chemical entity. The labile bond may be cleaved by a nucleophilic group, e.g. positioned on a scaffold

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WO 2005/026387

89

PCT/DK2004/000630

the subject of WO03078627A2, the content of which is incorporated herein in activator, i.e. a labile bond is formed between the oxygen atom connected to nucleophilic group, e.g. positioned on a scaffold, to transfer the chemical enifly to the scaffold, thus converting the remainder of the fragment into a leaving group of the reaction. When the chemical entity is connected to the actibond formed on the scaffold will an amide bond. The above building block is The 5-membered substituted N-hydroxysuccinimid (NHS) ring serves as an the NHS ring and the chemical entity. The labile bond may be cleaved by a vator through an carbonyl group and the recipient group is an amine, the their entirety by reference.

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Another building block which may form an amide bond is

R may be absent or NO2, CF3, halogen, preferably CI, Br, or I, and Z may be S or O. This type of building block is disclosed in WO03078626A2. The content of this patent application is incorporated herein in the entirety by refer-

group thereby transferring the chemical entity –(C=O)-CE' to said nucleo-A nucleophilic group can cleave the linkage between Z and the carbonyl phillc group. Building blocks transferring a chemical entity to a recipient reactive group forming a C=C bond

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recipient aldehylde group thereby forming a double bond between the carbon A building block as shown below are able to transfer the chemical entity to a

of the aldehyde and the chemical entity

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The above building block is disclosed in W003078445A2, the content of which being incorporated herein in the entirety by reference. 5

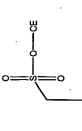
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Building blocks transferring a chemical entity to a recipient reactive group . forming a C-C bond

The below building block is able to transfer the chemical entity to a recipient group thereby forming a single bond between the receiving moiety, e.g. a scaffold, and the chemical entity. .20

PCT/DK2004/000630 5 WO 2005/026387

The above building block is disclosed in WO03078445A2, the content of which being incorporated herein in the entirety by reference. Another building block capable of transferring a chemical entity to a receiving reactive group forming a single bond is



atom, thereby forming a C-C bond between the chemical entity and the scafhetero atom, thereby forming a single bond between the chemical entity and fold. The above building block is disclose in WO03078446A2, the content of the hetero atom, or the receiving group may be an electronegative carbon The receiving group may be a nucleophile, such as a group comprising a which is incorporated herein by reference. 5

selected from a large arsenal of chemical structures. Examples of chemical The chemical entity attached to any of the above building blocks may be a entities are

kenyl, C₂-C₈ alkynyl, C₄-C₈ alkadienyl, C₃-C₇ cycloalkyl, C₃-C₇ cycloheteroal-H or entities selected among the group consisting of a C₁-C₆ alkyl, C₂-C₆ al-20

kylene-NR⁴C(O)OR⁸, C₁-C₂ alkylene-O-NR⁴2, C₁-C₂ alkylene-O-NR⁴C(O)R⁸, · kyl, aryl, and heteroaryl, said group being substituted with 0-3 R4, 0-3 R5 and 0-3 R³ or C₁-C₃ alkylene-NR⁴2, C₁-C₃ alkylene-NR⁴C(O)Rª, C₁-C₃ al-C₁-C₂ alkylene-O-NR⁴C(O)OR⁸ substituted with 0-3 R⁹.

of C₁-C₈ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyi, C₃-C₇ cycloalkyl, C₃-C₇ cyclohetwhere \mathbb{R}^4 is H or selected independently among the group consisting eroalkyl, aryl, heteroaryl, said group being substituted with 0-3 R9 and

-NHNHRs, -C(O)Rs, -SnRs, -B(ORs), -P(O)(ORs), or the group consisting of R5 is selected independently from -Ns. -CNO, -C(NOH)NH2, -NHOH, Cz-Ce alkenyl, Cz-Ce alkynyl, C₄-Ce alkadienyl said group being substituted

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where R^6 is selected independently from H, $C_1\text{-}C_6$ alkyl, $C_2\text{-}C_7$ cycloalkyl, aryl or C₁-C₈ alkylene-aryl substituted with 0-5 halogen atoms selected from -F, -Cl, -Br, and -I; and

R7 is independently selected from -NO2, -COOR6, -COR6, -CN, -OSIR³, -OR⁸ and -NR⁵₂.

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R⁸ is H, C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₃-C₇ cycloalkyl, агуl or С₁-С₈ alkylene-aryl substituted with 0-3 substituents independently selected from -F, -Cl, -NO2, -R³, -OR³, -SiR³3

R⁹ is =O, -F, -CI, -Br, -I, -CN, -NO₂, -OR⁶, -NR⁶₋₂, -NR⁶-C(O)R⁸, -NR $^{\rm c}$ -C(0)OR $^{\rm c}$, -S(0)R $^{\rm c}$, -S(0)2R $^{\rm c}$, -C(0)NR $^{\rm c}_2$ and

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Cross-link cleavage building blocks

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It may be advantageous to split the transfer of a chemical entity to a recipient cleavage step because each step can be optimized. A suitable building block reactive group into two separate steps, namely a cross-linking step and a for this two step process is illustrated below:

WO 2005/026387

PCT/DK2004/000630

pearing on a scaffold, thereby forming a cross-link. Subsequently, a cleavage initally, a reactive group appearing on the functional entlty precursor (abbreviated FEP) reacts with a recipient reactive group, e.g. a reactive group apis performed, usually by adding an aqueous oxidising agent such as 12, Br2, $\mathsf{Cl}_2,\mathsf{H}^\star$ or a Lewis acid. The cleavage results in a transfer of the group HZ-FEP- to the recipient moiety, such as a scaffold.

Z is O, S, NR⁴ In the above formula Q is N, CR1 9

C₁₋₈O-alkylene, C₁₋₈S-alkylene, NR¹-alkylene, C₁₋₈alkylene-S P is a valence bond, O, S, NR4, or a group $C_{57} \text{arylene,}$ option said group being substituted with 0-3 R4, 0-3 R5 and 0-3 R9 or C1-C3 alkylene-NR⁴2, C₁-C₃ alkylene-NR⁴C(O)R⁸, C₁-C₃ alkylene-NR⁴C(O)OR⁸ C₁-C₂ alkylene-O-NR⁴2, C₁-C₂ alkylene-O-NR⁴C(O)R⁸, C₁-C₂ alkylene-O-NR4C(O)OR8 substituted with 0-3 R9,

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B is a group comprising D-E-F, in which

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salkynylene, $C_{5.7}$ arylene, or $C_{5.7}$ heteroarylene, sald group optionally being D is a valence bond or a group C₁₋₆alkylene, C₁₋₆alkenylene, C₁. substituted with 1 to 4 group R11,

ealkylene, C_{1-s}alkenylene, C $_{1-8}$ alkynylene, C $_{5-7}$ arylene, or C $_{5-7}$ heteroarylene, E is, when present, a valence bond, O, S, NR4, or a group C,. said group optionally being substituted with 1 to 4 group R¹

F is, when present, a valence bond, O, S, or NR4,

A is a spacing group distancing the chemical structure from the complementing element, which may be a nucleic acid,

kylene-NR⁴2, C₁-C₃ alkylene-NR⁴C(O)R⁸, C₁-C₃ alkylene-NR⁴C(O)OR⁸, C₁-C₂ group consisting of H, C1-Ce alkyl, C2-Ce alkenyl, C2-Ce alkynyl, C4-Ce alkadi-R¹, R², and R³ are independent of each other selected among the enyl, C3-C7 cycloalkyl, C3-C7 cycloheteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 R⁴, 0-3 R⁵ and 0-3 R⁹ or C₁-C₃ alalkylene-O-NR²2, C₁-C₂ alkylene-O-NR⁴C(O)R⁸, C₁-C₂ alkylene-O-NR⁴C(O)OR⁸ substituted with 0-3 R⁹,

heteroalkyl, aryl, and heteroaryl, said group being substituted with 0-3 $\ensuremath{\text{R}}^4, 0\text{-}3$ FEP is a group selected among the group consisting of H, C₁-C₆ alkyl, Cz-Ce alkenyl, Cz-Ce alkynyl, C4-Ce alkadienyl, C3-C7 cycloalkyl, C3-C7 cyclo-R5 and 0-3 R9 or C1-C3 alkylene-NR42, C1-C3 alkylene-NR4C(O)R8, C1-C3 alkylene-NR*C(O)OR⁸, C₁-C₂ alkylene-O-NR*2, C₁-C₂ alkylene-O-NR*C(O)R⁸, C₁-C₂ alkylene-O-NR⁴C(O)OR⁸ substituted with 0-3 R⁹,

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of C₁-C₆ alkyl, C₂-C₆ alkenyl, C₂-C₆ alkynyl, C₃-C₇ cycloalkyl, C₃-C₇ cyclohetwhere R4 is H or selected independently among the group consisting eroalkyl, aryl, heteroaryl, said group being substituted with 0-3 R9 and

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-NHNHR⁸, -C(O)R⁸, -SnR³, -B(OR⁸)₂, -P(O)(OR⁶)₂ or the group consisting of R⁵ is selected independently from -N₃, -CNO, -C(NOH)NH₂, -NHOH, Cz-Ce alkenyl, Cz-Ce alkynyl, C4-Ce alkadienyl said group being substituted with 0-2 R7,

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where R⁶ is selected independently from H, C₁-C₆ alkyl, C₃C₇ cycloalkyl, aryl or C₁-C₆ alkylene-aryl substituted with 0-5 halogen atoms selected from -F, -Cl, -Br, and -I; and R⁷ is independently selected from -NO₂, -COOR⁶, -COR⁶, -CN, -OSIR⁵3, -OR⁸ and -NR⁶2.

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C₁-C₆ alkylene-aryl substituted with 0-3 substituents independently selected Re is H, C1-Ce alkyl, C2-Ce alkenyl, C2-Ce alkynyl, C3-C7 cycloalkyl, aryl or from -F, -CI, -NO2, -R3, -OR3, -SIR3

-NR⁶-C(0)OR⁶, -SR⁶, -S(0)R⁶, -S(0)₂R⁶, -COOR⁶, -C(0)NR⁶₂ and R⁸ is =0, -F, -CI, -Br, -I, -CN, -NO₂, -OR⁸, -NR⁸, -NR⁸-C(0)R⁸, -S(0)2NR⁶2. ജ

WO 2005/026387

95

PCT/DK2004/000630

CH2, and R1, R2, and R3 is H. The bond between the carbonyl group and Z is In a preferred embodiment Z is O or S, P is a valence bond, Q is CH, B is cleavable with aqueous I2.

Contacting between target and library

cleotide, may be referred to as the enrichment step or the selection step, as characteristics can include binding to a target, catalytically changing the tarjected under binding conditions to a target associated with a target oligonuget, chemically reacting with a target in a manner which alters/modifies the target or the functional activity of the target, and covalently attaching to the appropriate, and includes the screening of the library for display molecules The contacting step, by which the library of bifunctional molecules is subnaving predetermined desirable characteristics. Predetermined desirable target as in a suicide inhibitor.

ferred according to the present invention is to enrich molecules with respect molecules interact without immobilisation of the target. Displayed molecules played molecules in a certain aspect of the invention will be removed during In theory, display molecules of interest can be selected based on their propto binding affinity towards a target of interest. In a certain embodiment, the basic steps involve mixing the library of complexes with the immobilized tarthat bind to the target will be retained on this surface, while nonbinding disget of interest. The target can be attached to a column matrix or microtitre high-affinity interactions. In another embodiment, the target and displayed erties using either physical or physiological procedures. The method prewells with direct immobilization or by means of antibody binding or other

It may be considered advantageously to perform a chromatography step after plexes bound to the target can then be coupled to the target oligonucleotide. mobilized. After the coupling between the identifier oligonucleotide and the or instead of the washing step, notably in cases where the target is not im-

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a single or a series of wash steps. The identifier oligonucleotides of com-

96

target oligonucleotide, the coupled oligonucleotide may be recovered and optionally amplified before the decoding step.

A significant reduction in background binders may be obtained with increased washing volumes, repeating washing steps, higher detergent concentrations and prolonged incubation during washing. Thus, the more volume and number of steps used in the washing procedure together with more stringent conditions the more efficiently the non-binders and background binders will be removed. The right stringency in the washing step can also be used to remove low-affinity specific binders. However, the washing step will also remove wanted binders if too harsh conditions are used.

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은 . A blocking step, such as incubation of solid phase with skimmed milk proteins or other inert proteins and/or mild detergent such as Tween-20 and Triton X-100, may also be used to reduce the background. The washing conditions should be as stringent as possible to remove background binding but to retain specific binders that interact with the immobilized target. Generally, washing conditions are adjusted to maintain the desired affinity binders, e.g. binders in the micromolar, nanomolar, or pocomolar range.

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In traditional elution protocols, false positives due to suboptimal binding and washing conditions are difficult to circumvent and may require elaborate adjustments of experimental conditions. However, an enrichment of more than 100 to 1000 is rarely obtained. The present invention alleviates the problem with false positive being obtained because the non-specific binding complexes to a large extent remain in solution or attached to the reaction chamber such that the indentifier oligonucleotide of non-binding complexes will be in a low concentration compared to the identifier oligonucleotides of binding complexes relative to the concentration of the target oligonucleotide.

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The target can be any compound of interest. E.g. the target can be a protein, peptide, carbohydrate, polysaccharide, glycoprotein, hormone, receptor, an-

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WO 2005/026387

26

PCT/DK2004/000630

tigen, antibody, virus, substrate, metabolite, transition state analogue, cofactor, inhibitor, drug, dye, nutrient, growth factor, cell, tissue, etc. without limitation. Sultable targets include, but are not limited to, angiotensin converting enzyme, renin, cyclooxygenase, 5-lipoxygenase, IIL- 10 converting enzyme, cytokine recentors. PDGF recentor type II nosine manaphase, debydox

cytokine receptors, PDGF receptor, type II Inosine monophosphate dehydrogenase, β-lactamases, integrin, proteases like factor VIIa, kinases like Bcr-Abl/Her, phosphotases like PTP-18, and fungal cytochrome P-450. Targets can include, but are not limited to, bradykinin, neutrophil elastase, the HIV proteins, including *tat, rav, gag, int*, RT, nucleocapsid etc., VEGF, bFGF, TGFB, KGF, PDGF, GPCR, thrombin, substance P, tgE, sPLA2, red blood cells, glioblastomas, fibrin clots, PBMCs, hCG, lectins, selectins, cytokines, ICP4, complement proteins, etc.

Certain targets comprise one or more discrete binding domains. Proteins that contain these domains are involved in a variety of processes, such as cellular transporters, cholesterol movement, signal transduction and signaling functions which are involved in development and neurotransmission. See Herz, Lipóprotein receptors: beacons to neurons?, (2001) <u>Trends in Neurosciences</u> 24(4):193-195; Goldstein and Brown,The Cholesterol Quartet, (2001) <u>Sci.</u>

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ence 292:1310-1312. The function of a discrete binding domain is often specific but it also contributes to the overall activity of the protein or polypeptide. For example, the LDL-receptor class A domain (also referred to as a class A module, a complement type repeat or an A-domain) is involved in ligand binding while the gamma-carboxyglumatic acid (Gla) domain which is found in the vitamin-K-dependent blood coagulation proteins is involved in highaffinity binding to phosphollpid membranes. Other discrete binding domains include, e.g., the epidermal growth factor (EGF)-like domain in tissue-type plasminogen activator which mediates binding to liver cells and thereby regulates the clearance of this fibrinolytic enzyme from the circulation and the cytoplasmic tail of the LDL-receptor which is involved in receptormediated en-

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tein Receptor Family, (1999) Annu. Rev. Nutr. 19:141-72. For example, some tion by lysosomes. See Hussain et al., The Mammalian Low-Density Lipopromains, as discussed above. These proteins are often called mosaic proteins. ogy domain containing YWTD repeats; 6) single membrane-spanning region; precursor-like repeats, a transmembrane domain and a cytoplasmic domain. protein E receptor 2, LDLR-related protein (LRP) and megalin. Family memcalcium for ligand binding; 4) recognition of receptor-associated protein and sent invention offers the possibility of identifying two or more ligands against tural domains: the cysteine rich A-domain repeats, epidermal growth factor lors; 2) recognize extracellular ligands; and 3) internalize them for degrada-For example, members of the LDL-receptor family contain four major structracellular ligand binding consisting of A-domain repeats; 3) requirement of members include very-low-density lipoprotein receptors (VLDL-R), apolipoapolipoprotein (apo) E; 5), epidermal growth factor (EGF) precursor homoland 7) receptor-mediated endocytosis of various ligands. See Hussain, supra. Yet, the members bind several structurally dissimilar ligands. The pre-The LDL-receptor family includes members that: 1) are cell-surface recep-Individual target proteins can possess one or more discrete monomer dobers have the following characteristics: 1) cell-surface expression; 2) exthe same target as discussed elsewhere herein.

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In some aspects of the invention, a dimer compound binding with two targets normally interacting in a biological context is identified using the methods of the present invention. Examples of such targets are factor Xa and factor Vila. The method includes separately screening a library of bifunctional complexes and identifying suitable display molecules that binds to both targets. The two display molecules pools are then linked together. In the event m display molecules are identified having an affinity above a certain threshold (i.e. a low Kd) to a first target and n display molecules having an affinity above another or the same threshold, a dimer compound array of n times m molecules are formed. The dimer compound are subsequently screened for the ability to bind to the first and the second target molecule, thereby identifying a dimer or

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WO 2005/026387 PCT/DK2004/000630

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a range of dimers that specifically bind to the first and the second target molecule with a certain affinity.

A target can also be a surface of a non-biological origin, such as a polymer surface or a metal surface. The method of the invention may then be used to identify sultable coatings for such surfaces.

In a preferred embodiment, the desirable display molecule acts on the target without any interaction between the nucleic acid attached to the desirable encoded molecule and the target. In one embodiment, the bound complex-target aggregate can be partitioned from unbound complexes prior to or subsequent to the coupling step by a number of methods. The methods include nitrocellulose filter binding, column chromatography, filtration, affinity chromatography, centrifugation, and other well known methods. A preferred method is size-exclusion chromatography.

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Briefly, the library of complexes is subjected to the target, which may include contact between the library and a column onto which the target is immobilised. Identifier oligonucleotides associated with undesirable display molecules, i.e. display molecules not bound to the target under the stringency conditions used, will pass through the column. Additional undesirable display molecules (e.g. display molecules which cross-react with other targets) may be removed by counter-selection methods. Desirable complexes are bound to the column. The target may be immobilized in a number of ways. In one embodiment, the target is immobilized through a cleavable physical link, such as one more chemical bonds. Following the interaction of the display molecule and the target, the respective oligonucleotides are coupled. The aggregate of the target and the complex may then be subjected to a size exclusion chromatography to separate the aggregate from the rest of the compounds in

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the cleavable linker that attached the target to the solid support. Subsequent cleaved to separate the Identifier oligonucleotides of complexes having affinty towards the targets. Just to mention a single type of orthogonal cleavable inkages, one could attached to target to the solid support through a linkage play molecule and the identifier oligonucleotide may be selected as a photocleavable linkage. More specifically, the former linkage may be a disulphide The complex may be provided with a cleavable linker at a position between immobilized the cleavable linker of the complex is preferable orthogonal to that can be cleaved by a chemical agent, and the linker separating the disbond that can be cleaved by a suitable reducing agent like DTT (dithiothreithe display molecule and the identifier oligonucleotide. When the target is to the optional size exclusion chromatography, this cleavable linker is ol) and the latter linkage may be an o-nitrophenyl group.

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method. In one embodiment, the coupling products can be fractionated by a There are other partitioning and screening processes which are compatible number of common methods and then each fraction is assayed for activity. with this invention that are known to one of ordinary skill in the art. Such known process may be used in combination with the present inventive The fractionization methods can include size, pH, hydrophobicity, etc. 5

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selection), followed by positive selection with the desired target. As an exam-Inherent in the present method is the selection of encoded molecules on the ing the selection process by first extracting complexes which are capable of cules with a desired function and specificity. Specificity can be required durple, inhibitors of fungal cytochrome P-450 are known to cross-react to some extent with mammalian cytochrome P-450 (resulting in serious side effects). Highly specific inhibitors of the fungal cytochrome could be selected from a basis of a desired function; this can be extended to the selection of moleibrary by first removing those complexes capable of interacting with the interacting with a non-desired "target" (negative selection, or counter-

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WO 2005/026387

PCT/DK2004/000630

mammalian cytochrome, followed by retention of the remaining products which are capable of interacting with the fungal cytochrome.

This can be done manually or using a more automatic system such as robotic inked to an oligonucleotide that Identifies the structure of the said compound. identify compounds that specifically interact with a certain protein kinase. The ent molecular targets. A small library of preferred compounds can be directly use of the present proximity selection procedure will in this instance generate The present invention can be used to identify compounds that bind to differequipments. These tagged compounds can then be mixed with one or more cules. The above method can also be used for other target classes such as study the selectivity and specificity and to design sub-libraries with potential target molecules to select for compound and target pairs that bind to each other. For example, a library of compounds designed to bind preferably to protein kinases could be mixed with a library of various protein kinases to compound are match simultaneously against different related target molean extensive structure activity relationship (SAR) where different binding proteases, phosphatases, GPCRs, nuclear receptors and corresponding compound libraries. The information for these selections can be used to binding compounds. S 9 रु 2

binding constant can be captured if the target concentration is higher than the in a prior art selection where the target is immobilized to a surface, for exambetween the molecular target and the binding molecules. The invention is not tocols. The amount of captured binding molecules can be varied using a suitmolecules are free in the solution and removed in future washing steps. The able concentration of the target. Most of the binding molecules with a certain dependent on washing- or separation-step as is most prior art selection prople in a well or a bead, the effective concentration of the target will be high present invention allows selection in solution at true equilibrium conditions locally on the surface but infinitively low in the solution. This restriction will result in low recovery of binding molecules because most of the binding

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binding constant of the binding molecules. Also the concentration of the target can be adjusted to capture binders with a certain binding constant. A high target concentration will also increase the likelihood of selecting specific binders that are present in low copy number. By using high concentration of target, the solution selection can also be used to identify binders with low binding affinity. This is especially important with screening for small fragments which normally possess low affinity but still holds important structural information that can be used in second generation library design.

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In a certain embodiment, a binding platform may be constructed that can be used for almost any target. The binding platform should preferably be small enough to only allow association of a few or a single target molecule. This to ensure a solution based selection procedure with adjustable target concentration. The binding platform is primarily composed of two components; a small surface allowing association of the target molecule, and an association area/site for the target oligonucleotide. This binding platform may be designed to mediate the association of the target and target oligonucleotide to allow proximity selection in solution.

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20 · Cleavable linkers

A cleavable linker may be positioned between the target and a solid support, between the potential drug candidate and the identifier oligonucleotide, between the molecular target and the target oligonucleotide or any other position that can provide for a separation of the identifier oligonucleotide from successful complexes from non-specific binding complexes. The cleavable linker may be selectively cleavable, i.e. conditions may selected that only cleave that particular linker.

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The cleavable linkers may be selected from a large plethora of chemical structures. Examples of linkers includes, but are not limited to, linkers having an enzymatic cleavage site, linkers comprising a chemical degradable component, and linkers cleavable by electromagnetic radiation, such as light.

WO 2005/026387

103

PCT/DK2004/000630

Examples of linkers cleavable by electromagnetic radiation (light)

o-nitrobenzyl

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R3

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o-nitrobenzyl in exo position

For more details see Holmes CP. J. Org. Chem. 1997, 62, 2370-2380

3-nitrophenyloxy

For more details see Rajasekharan Pillai, V. N. Synthesis. 1980, 1-26

Dansyl derivatives:

For more details see Rajasekharan Pillai, V. N. Synthesis. 1980, 1-26

Coumarin derivatives :

For more details see R. O. Schoenleber, B. Giese. Synlett 2003, 501-504

gonucleotide, respectively. Alternatively, R1 and R2 can be either of the target R¹ and R² can be either of the potential drug candidate and the identifier olior a solid support, respectively.

R3 = H or OCH3

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WO 2005/026387

PCT/DK2004/000630

If X is O then the product will be a carboxylic acid If X is NH the product will be a carboxamide

catalog # 10-4913-90) which can be introduced in an oligonucleotide during synthesis and cleaved by subjecting the sample in water to UV light (~ 300-One specific example is the PC Spacer Phosphoramidite (Glen research 350 nm) for 30 seconds to 1 minute.

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DMT = 4,4'-Dimethoxytrityl CNEt = Cyanoethyl iPr = Isopropyl

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complexes at a position between the indentifier and the potential drug candi-The above PC spacer phosphoamidite is suitable incorporated in a library of date. The spacer may be cleaved according to the following reaction.

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cule, respectively. In a preferred aspect R2 is an oligonucleotide identifier and phate group is generated allowing for further biological reactions. As an ex-R¹ and R² can be either of the encoded molecule and the identifying moleample, the phosphate group may be positioned in the 5'end of an oligonuthe R1 is the potential drug candidate. When the linker is cleaved a phoscleotide allowing for an enzymatic ligation process to take place.

Examples of linkers cleavable by chemical agents:

Ester linkers can be cleaved by nucleophilic attack using e.g. hydroxide ions. In practice this can be accomplished by subjecting the target-ligand complex to a base for a short period.

R¹ and R² can be the either of be the potential drug candidate or the identifier oligonucleotide, respectively. R⁴6 can be any of the following: H, CN, F, NO₂, SO₂NR₂.

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Disulfide linkers can efficiently be cleaved / reduced by Tris (2-carboxyethyl) phosphine (TCEP). TCEP selectively and completely reduces even the most stable water-soluble alkyl disulfides over a wide pH range. These reductions frequently required less than 5 minutes at room temperature. TCEP is a nonvolatile and odorless reductant and unlike most other reducing agents, it is resistant to air oxidation. Trialkylphosphines such as TCEP are stable in aqueous solution, selectively reduce disulfide bonds, and are essentially unreactive toward other functional groups commonly found in proteins.

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More details on the reduction of disulfide bonds can be found in Kirley,

T.L.(1989), Reduction and fluorescent labeling of cyst(e)ine-containing pro-

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WO 2005/026387

107

PCT/DK2004/000630

teins for subsequent structural analysis, Anal. Biochem. 180, 231 and Levison, M.E., et al. (1969), Reduction of biological substances by water-soluble phosphines: Gamma-globulin. Experentia 25, 126-127.

5 Linkers cleavable by enzymes

The linker connecting the potential drug candidate with the identifier oligonucleotide or the solid support and the target can include a peptide region that allows a specific cleavage using a protease. This is a well-known strategy in molecular biology. Site-specific proteases and their cognate target amino acid sequences are often used to remove the fusion protein tags that facilitate enhanced expression, solubility, secretion or purification of the fusion protein.

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Various proteases can be used to accomplish a specific cleavage. The specificity is especially important when the cleavage site is presented together with other sequences such as for example the fusion proteins. Various conditions have been optimized in order to enhance the cleavage efficiency and control the specificity. These conditions are available and know in the art.

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Enterokinase is one example of an enzyme (serine protease) that cut a specific amino acid sequence. Enterokinase recognition site Is Asp-Asp-Asp-Asp-Lys (DDDDK), and it cleaves C-terminally of Lys. Purified recombinant Enterokinase is commercially available and is highly active over wide ranges in pH (pH 4.5-9.5) and temperature (4-45°C).

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The nuclear inclusion protease from tobacco etch virus (TEV) is another commercially available and well-characterized proteases that can be used to cut at a specific amino acid sequence. TEV protease cleaves the sequence Glu-Asn-Leu-Tyr-Phe-Gln-Gly/Ser (ENLYFQG/S) between Gln-Gly or Gln-Ser with high specificity.

Another well-known protease is thrombin that specifically cleaves the sequence Leu-Val-Pro-Arg-Gly-Ser (LVPAGS) between Arg-Gly. Thrombin has

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quences can also be used for thrombin cleavage; these sequences are more or less specific and more or less efficiently cleaved by thrombin. Thrombin is a highly active protease and various reaction conditions are known to the also been used for cleavage of recombinant fusion proteins. Other se-

Activated coagulation factor FX (FXa) is also known to be a specific and useful protease. This enzyme deaves C-terminal of Arg at the sequence lle-Glu-Gly-Arg (IEGR). FXa is frequently used to cut between fusion proteins when producing proteins with recombinant technology. Other recognition sequences can also be used for FXa.

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Other types of proteolytic enzymes can also be used that recognize specific acid sequences in an un-specific manner can also be used if only the linker amino acid sequences. in addition, proteolytic enzymes that cleave amino contains an amino acid sequence in the complex molecule.

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Other type of molecules such as ribozymes, catalytically active antibodies, or ipases can also be used. The only prerequisite is that the catalytically active the linker, that connects the encoding region and the displayed molecule or, molecule can cleave the specific structure used as the linker, or as a part of in the altemative the solid support and the target.

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quence is close to the nucleotide sequence length. Purified recombinant Eco A variety of endonucleases are available that recognize and cleave a double donuclease Eco RI is an example of a nuclease that efficiently cuts a nucleotions. As an example the Eco RI is working in in various protocols as indicted tide sequence linker comprising the sequence GAATTC also when this se-RI is commercially available and is highly active in a range of buffer condistranded nucleic acid having a specific sequence of nucleotides. The enbelow (NEBuffer is available from New England Biolabs):

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WO 2005/026387

PCT/DK2004/000630

NEBuffer 1: [10 mM Bis Tris Propane-HCI, 10 mM MgCl2, 1 mM dithiothreftol (pH 7.0 at 25°C)],

NEBuffer 2 : [50 mM NaCl, 10 mM Tris-HCl, 10 mM MgCl2, 1 mM dithiothreltol (pH 7.9 at 25°C)],

NEBuffer 3: [100 mM NaCl, 50 mM Tris-HCl, 10 mM MgCl2, 1 mM dithlothreitol (pH 7.9 at 25°C)], NEBuffer 4 : [50 mM potassium acetate, 20 mM Tris-acetate, 10 mM magnesium acetate, 1 mM dithiothreitol (pH 7.9 at 25°C)] Extension buffer: mM KCI, 20 mM Tris-HCI(Ph 8.8 at 25o C), 10 mM (NH4)2 SO4, 2 mM MgSO 4 and 0.1% Triton X-100, and 200 µM dNTPs. 2

Determining the identifier oligonucleotide sequence

entities as well as the point in time they have been incorporated in the display cule(s) and 7 or optionally the molecular target(s). In a certain embodiment of play molecules that binds to the target are identified. The synthesis method molecule can be deduced from the identifier oligonucleotide. It may be suffi-The nucleotide sequence of the identifier sequence present in the coupled the invention, chemical entities that participated in the formation of the disof the display molecule may be established if information on the chemical product is determined to identify the identity of the binding display mole-5

hindrance on the scaffold molecule or the chemical entity to be transferred. In entities that have participated in the display molecule to deduce the full molecient to obtain information on the chemical structure of the various chemical use of different kinds of attachment chemistries may ensure that a chemical cule due to structural constraints during the formation. As an example, the scaffold. Another kind of chemical constrains may be present due to steric identifier oligonucleotide sequence that enable the identification of each of entity on a building block can only be transferred to a single position on a general however, it is preferred that information can be inferred from the 25 8

the chemical entities that have participated in the formation of the encoded

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molecule along with the point in time in the synthesis history the chemical entities have been incorporated in the (nascent) display molecule.

Although conventional DNA sequencing methods are readily available and useful for this determination, the amount and quality of isolated bifunctional molecule may require additional manipulations prior to a sequencing reaction.

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Where the amount is low, it is preferred to increase the amount of the coupled oligonucleotide sequence by polymerase chain reaction (PCR) using PCR primers directed to primer binding sites present in the identifier oligonucleotide sequence.

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In one embodiment, the different coupled oligonucleotide sequences are cloned into separate sequencing vectors prior to determining their sequence by DNA sequencing methods. This is typically accomplished by amplifying the different coupled oligonucleotide sequences by PCR and then using a unique restriction endonuclease sites on the amplified product to directionally clone the amplified fragments into sequencing vectors. The cloning and sequencing of the amplified fragments then is a routine procedure that can be carried out by any of a number of molecular biological methods known in the art.

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Alternatively, the bifunctional complex or the PCR amplified identifier oil-gonucleotide sequence can be analysed in a microarray. The array may be designed to analyse the presence of a single codon or multiple codons in a identifier oligonucleotide sequence.

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In still another approach, the coupled oligonucleotide product is analysed by QPCR. Preferably, the QPCR affords information as to the chemical moieties that has participated in the formation of the display molecules and optionally the identity of the target. The QPCR approach also allows a direct investiga-

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WO 2005/026387

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PCT/DK2004/000630

tion of the enrichment factor if two samples are analysed in parallel, one with target and the other with the target plus library. The difference in signal from these to samples will illustrate how much coupling that is due to the target mediated coupling compared to the background coupling. Various conditions can be investigated to obtain the most optimal selection procedure before the sequences are analysed to identify the precise structures of the binding molecules.

Synthesis of nucleic acids

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Oligonucleotides can be synthesized by a variety of chemistries as Is well known in the art. For synthesis of an oligonucleotide on a substrate in the direction of 3' to 5', a free hydroxy terminus is required that can be conveniently blocked and deblocked as needed. A preferred hydroxy terminus blocking group is a dimexothytrityl ether (DMT). DMT blocked termini are first deblocked, such as by treatment with 3% dichloroacetic acid in dichloromethane (DCM) as is well known for oligonucleotide synthesis, to form a free hydroxy terminus.

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Nucleotides in precursor form for addition to a free hydroxy terminus in the direction of 3' to 5' require a phosphoramidate moiety having an aminodiisopropyl side chain at the 3' terminus of a nucleotide. In addition, the free hydroxy of the phosphoramidate is blocked with a cyanoethyl ester (OCNET), and the 5' terminus is blocked with a DMT ether. The addition of a 5' DMT-, 3' OCNET-blocked phosphoramidate nucleotide to a free hydroxyl requires tetrazole in acetonitrile followed by iodine oxidation and capping of unreacted hydroxyls with acetic anhydride, as is well known for oligonucleotide synthesis. The resulting product contains an added nucleotide residue with a DMT blocked 5' terminus, ready for deblocking and addition of a subsequent blocked nucleotide as before.

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For synthesis of an oligonucleotide in the direction of 5' to 3', a free hydroxy terminus on the linker is required as before. However, the blocked nucleotide

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to be added has the blocking chemistries reversed on its 5' and 3' termini to facilitate addition in the opposite orientation. A nucleotide with a free 3' hydroxyl and 5' DMT ether is first blocked at the 3' hydroxy terminus by reaction with TBS-CI in imidazole to form a TBS ester at the 3' terminus. Then the

DMT-blocked 5' terminus is deblocked with DCA in DCM as before to form a free 5' hydroxy terminus. The reagent (N.N-diisopropylamino)(cyanoethyl) phosphonamidic chloride having an aminodiisopropyl group and an OCNET ester is reacted in tetrahydrofuran (THF) with the 5' deblocked nucleotide to form the aminodiisopropyl-, OCNET-blocked phosphonamidate group on the 5' terminus. Thereafter the 3' TBS ester is removed with tetrabutylammonium fluoride (TBAF) in DCM to form a nucleotide with the phosphonamidateblocked 5' terminus and a free 3' hydroxy terminus. Reaction in base with DMT-Cl adds a DMT ether blocking group to the 3' hydroxy terminus.

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The addition of the 3' DMT-, 5' OCNET-blocked phosphonamidated nucleotide to a linker substrate having a free hydroxy terminus then proceeds using the previous tetrazole reaction, as is well known for oligonucleotide polymerization. The resulting product contains an added nucleotide residue with a DMT-blocked 3' terminus, ready for deblocking with DCA in DCM and the addition of a subsequent blocked nucleotide as before.

Brief Description of the Figures

Fig. 1 discloses an embodiment for proximity-dependent selection,

Fig. 2 discloses different approaches for accomplishing coupling,

25 Fig. 3 discloses four different approaches for producing a coupling

Fig. 4 discloses a library versus library screening,
Fig. 5 discloses an embodiment in which a target oligonucleotide association
is performed in solution.

Fig. 6 discloses a target oligonucleotide association on cell surface.

Fig. 7 discloses a target with multiple binding sites which may associate with members of the bifunctional molecule library.

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plexes in solution to avoid unspecific ligation.

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WO 2005/026387

PCT/DK2004/000630

Fig. 8 discloses a target with one binding site for association with a pair of displayed molecules.

Fig. 9 discloses a 2nd generation-library driven proximity selection. Fig. 10 discloses multiple targets for simultaneously subjected to a library of

5 complexes.

Detailed Disclosure of the Figures

Fig. 1 outlines an embodiment for a proximity-dependent selection (PDS). The molecular target is linked to a target oligonucleotide, which in some embodiment may be unique for the target molecule. This target sequence comes in close proximity with a specific identifier oligonucleotide when the displayed molecule of a bifunctional complex binds to the target molecule. This proximity will promote the coupling between the bifunctional complex molecules that bind to the target compare to bifunctional complex in solution.

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Thus, there will be a selection for coupling products that contain display molecules that possess affinity for the target molecule. The final ligation product is amplified using two primers that only amplify ligated products.

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In a first step, the target associate with a target oligonucleotide is mixed with a library of complexes, in which each complex comprises a display molecule attached to an identifier oligonucleotide. The display molecules are then incubated with the target. The display molecules which have an affinity towards the molecular target will bind, while the complexes not having affinity will remain in solution. Subsequent to the incubation, a connector oligonucleotide is added. The connector oligo nucleotide comprises parts that hybridise to sequences near the ends of the target and the identifier oligonucleotides, respectively. Subsequent to the addition of the connector oligonucleotide, a ligation is effected by chemical or enzymatic means. Preferably a ligase is used to ligate the target and the identifier oligonucleotides together. The connector oligonucleotide is generally added in excess to saturate the comnector oligonucleotide is generally added in excess to saturate the com-

114

After the ligation, the ligation product is amplified by PCR. Thus, a forward primer is annealed to the ligation product at the 3' end thereof and extended using a polymerase. The transcribed product comprises a site to which a second (or reverse) primer can anneal so as to provide for an extension of the second primer using the transcribed product as template. Using forward and reverse primers as indicated above together with a polymerase and suitable substrates produces amplicons, which comprises information about the display molecule as well as the molecular target. The figated product can be introduced into a host organism using a suitable vector. The host vector may be allowed to form colonies and the colonies can be sequenced to establish the identity of the display molecule.

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Fig. 2 shows various options to perform coupling between the target oligonucleotide and the identifier oligonucleotide. A. The ligation is promoted using a connector oligonucleotide that anneals both to the target oligonucleotide and the identifier oligonucleotide. The connector oligonucleotide is designed such that the ends of the identifier oligonucleotide and the target oligonucleotides are abutted. A ligase is subsequently allowed to ligate the ends together. B. A connector oligonucleotide is used to promote fill in of a gap using a polymerase and finally ligation using a ligase. C. The distal end of the target oligonucleotide overlaps the distal end of the identifier oligonucleotide, which allows a polymerase to extend the target oligonucleotide as well as the identifier oligonucleotide thereby forming a double stranded product. D. Bluntended ligation of single-stranded or double stranded DNA using a suitable enzyme like T4 DNA ligase.

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Fig. 3 shows various methods for preparing a coupling area on an existing bifunctional complex. Conjugates between molecular targets associated with an oligonucleotide and complexes comprising a display molecule and an identifier oligonucleotide can be modified to allow a ligase to couple the oligonucleotides together. A. The identifier oligonucleotide is extended with a primer with an overhang that creates the coupling area. The extension is

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WO 2005/026387

PCT/DK2004/000630

suitably conducted before the selection process to obtain the benefit of a double stranded nucleotide sequence. A target oligonucleotide can be ligated to the blunt end of the extended primer or a connector oligonucleotide can be used to connect the target oligonucleotide and the extended primer prior to ligation with a suitable ligase. B. The identifier oligonucleotide is annealed to a primer that blinds internally. The primer is subsequently extended, sultably before the selection process. The extension forms a coupling area directly on the identifier oligonucleotide, which allows a target oligonucleotide to be annealed and ligated. C. The first step is identical to the procedure as describe in B but the target sequence has a free 5'-end that allow ligation to the 3'-end of the identifier oligonucleotide. A blunt ended single stranded ligation can be

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in B but the target sequence has a free 5'-end that allow ligation to the 3'-end of the identifier oligonucleotide. A blunt ended single stranded ligation can be performed. Alternatively, this variation can be performed using a connector oligonucleotide and subsequent ligation. D. A primer is annealed to a identifier oligonucleotide and extended to produce a double-stranded DNA which is subsequently cut with an enzyme (e.g. restriction enzyme) to produce a single-stranded DNA portion that can be used as handle in the coupling process.

Fig. 4 shows a library versus library selection method. Different targets specifically encoded by the attached target oligonucleotides are mixed with a library of bifunctional complexes. The displayed molecules will bind to specific targets and promote the ligation through the proximity effect. This ligation will connect the target oligonucleotides with oligonucleotides that encodes for specific displayed molecules. The ligated oligonucleotides can be amplified and determined by sequencing procedures well known in the art. The ligated sequences will reveal which display molecules that bind to which target.

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Fig. 5 discloses *inter alia* the association of the target oligonucleotide to the target. One way of associating the target oligonucleotide with the target molecule is to link the oligonucleotide through a tag introduced on the target molecule. The tag can be attached before the target is produced (e.g. a short

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amino acid sequence such as HIS-tag of FLAG-tag) or be modified after the target is produced. The target sequence can then be associated through the tag using a tag-binding molecule such as an antibody or other type of molecules that binds to the tag.

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Fig. 6 discloses target oligonucleotide association on a cell surface. Specific receptors can be engineered to express a specific tag on the cell surface. Different tags can be used such as HIS- or FLAG-tags or other types of tags that become bound with the receptor. The tag will only be displayed on the cell surface together with the specific receptor. The target oligonucleotide is then associated with the receptor target using a mediator molecule that carries the target oligonucleotide and binds to the tag. A mediator molecule could be an antibody that binds to the tag (e.g. anti-HIS or anti-FLAG antibodies) that is associated with the target oligonucleotide. This procedure will specifically associate the target oligonucleotide with a receptor target on the cell surface which will promote a ligation between oligonucleotides of the binding displayed molecules and the target oligonucleotide.

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Fig. 7 shows a target molecule with several sites for binding of ligands. The target is subjected to a library of complexes of bifunctional molecules. Display molecules of the complexes binds to the discrete sites of the molecular target thus promoting a high local concentration of the ends of the oligonucleotides which have bound to the target. Subsequently a connector oligonucleotide is added to adjoin the distal ends of the oligonucleotides together. Usually, the connector oligonucleotide is added in excess to saturate the ends of the identifier oligonucleotides free in the solution. The ends of the oligonucleotides kept together by the connector oligonucleotide are ligated together forming a coupled product. The coupled product is amplified by PCR using primers annealing to each end of the coupled product. The amplified coupled product is decoded to identify the display molecules which have bound to the in the target. In a step not shown on the figure, the two binding display molecules are coupled together via a suitable linker to form a ligand

32

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WO 2005/026387

117

PCT/DK2004/000630

which binds to two sites of the target. Suitable, the dimer comprising the two revealed display molecules and the linker is synthesised by organic synthesis.

Libraries of bifunctional complexes can also be screened against each other using the present invention. Such an embodiment allows the detecting of pairs of displayed molecules that bind to the same target at different or the same binding site or pair of displayed molecules that bind to different targets.

The power of the screening libraries in the above fashion is indicated by the fact that a library of e.g. 10⁴ different displayed molecules generates a total combination of display molecules of 10⁸ when pair of binders are searched

external reactant so as to form a single molecule. After the binding interaction tially, the target is mixed with the library of bifunctional complexes under conget oligonucleotide. Subsequently to or simultaneously with the binding of the covalent linkage between the display molecules. In another embodiment, the complex associates with the target to form the target associated with the tarfirst display molecule, a second complex binds to the same site of the target. The display molecules may or may not be reacted with each other to form a comprises display molecules that binds to the target are joint together. In an two display molecules are connected via a suitable linker or reacted with an to saturate ends of identifier oligonucleotides which are not part of a binding ditions which promote a binding interaction to take place. A first bifunctional of the library of complexes with the target, the ends of the complexes which polynucleotide. The connector polynucleotide is preferably added in excess complex. After the hybridisation event between the ends of the identifier oll-Fig. 8 discloses a library of bifunctional complexes which is presented to a target having a site possible to be occupied by two dlsplay molecules. Inigonucleotides and the connector oligonucleotide a ligation is conducted. aspect of the invention, the ends are joined together using a connector Suitably the ligation is performed by a ligase to form a coupled product, 5 ೫ 25 ဓ္က

formation of the display molecules which have participated in the binding incoupled product is amplified by PCR to form PCR amplicons comprising inwhich can be used as a template by a polymerase. After the ligation, the eraction. Fig. 9 discloses a two (or more) step identification method. In a first step the generate the second generation library. Initially, a library of complexes are is with further components and/or the low binding display molecule is added or on the drawing with a display molecule only having a partial fit in the binding library prepared upon the knowledge harvested in the first library is used to presented to a target having a binding site. In the library, display molecules having a binding affinity above a certain threshold is not present, illustrated nents used in the synthesis of the low binding display molecule are shuffled method as disclosed in figure 1 is conducted and in the second step a new site of the target. In the synthesis of the second generation library compo-

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cule has been successful display molecules are generated which binds with a ing on the natural translation system a deletion, alteration or addition of nucleic acid can be performed. The second generation library is presented to the a target again. In the event the alteration of the initial low binding molehigher affinity towards the target. ឧ

chemical entities can be conducted. For systems for complex generation rely-

subtracted a structural unit. As an example, a further round of addition of

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attachment can be artificial, i.e. the association between target 1 and target 2 association between target 1 and target 2 occur in a biological context or the which ensure a linkage. The association of the targets may also be obtained mixing with the library of complexes. The attachment can be natural, i.e. the between the targets may be obtained by any chemical or enzymatic means naving two distinct targets or monomer domains. In an embodiment, one of by expressing target 1 and target 2 as a fusion protein, i.e. a single protein Fig. 10 discloses two targets which are attached to each other prior to the is obtained by a chemical synthesis. In the latter instance, the association 23 ဓ

WO 2005/026387

PCT/DK2004/000630

event the library is spiked with complexes having a ligand against the capturthe targets in the fusion protein is a capturing protein, like streptavidin. In the protein and a member of the library. The further functionality, i.e. target 2, of ing protein, like blotin, it is feasible to form a connection between the fusion the fusion protein may be then be subjected to a screening process to find binder from the library.

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During the mixing step, the two attached targets are contacted with the library comprises genetic information which encodes both the display molecules that suitable binding display molecules, two ends of the binding complexes is pocomplex comprising a compounds known to bind to the one of the targets in order to find sultable binders against another target. If the library comprises sures that the ends are kept close together when a ligase is allowed to persitioned in close proximity. The addition of a connector oligonucleotide enof complexes under binding conditions. The library may be spiked with a form the action of ligating the ends together. The resulting PCR product have participated in the binding interaction with target 1 and target 2.

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EXAMPLES illustrating the second aspect of the present invention Example 1:

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Oligonucleotide sequences

Target Sequence (ES-1) 22

5'-X-TAGTC GATGT AGCTA GCTAG TGCGC CAATG CCTTA TCAGC

5'-GATCG ATGAC TGACG CCGGT AAATCTACCGTCTAAGCTG-Y-3' Identifier Sequence (IS-1) (extension part)

Underlined sequence is reverse primer binding site

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Control Identifier Sequence (CIS-1) (extension part)

120

5'-CANCG ANGAC TGACG CCGGT gacgt cgtag atatc gatgc AAATCTACCGTCTAAGCTG-Z-3'

Underlined sequence is reverse primer

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Connector Sequence (CS-1) 5' - AAAAAGAAATAGTCG-CTAGCTACTGTTTT Primers (Forward PR-1 and reverse PR-2)

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PR-1: 5'-TAGTC GATGT AGCTA GCTAG PR-2: 5'-CAGCT TAGAC GGTAG ATTT Target labelling with oligonucleotide seguence.

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The target molecule (streptavidin) is modified with an oligonucleotide sequence using a terminus modifier that allow direct coupling to the target(s) molecule. The oligonucleotide sequence ES-1 is synthesised with the 5'Thiol-Modifier (Glen Research, #10-1926-90) to produce a oligonucleotide that can be coupled to the target(s) through the 5'-end (designated X in the

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-Thiol-Modifier C6

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The covalent attachment of the oligonucleotide at the target is carried out with the aid of the heterobispecific crosslinker Sulfosuccinimidyl-4-(N-maleimidomethyl)cyclohexane-1-carboxylate (sSMMCC). The ɛ-amino groups of lysine side chains of the target(s) are first derivatized with sSMCC cross linker to provide a maleimide functionalify, which subsequently is reacted with the thiolated oligonucleotide.

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WO 2005/026387

PCT/DK2004/000630

121

Protecol: Dissolve approx. 2 mg of sSMCC in 60 µl of DMF. Add the sSMCC solution to 200 µl of a 100 µM solution of streptavidin in PBS buffer pH 7.3 and incubate in the dark at room temperature for about 1 hour. The excess sSMCC is removed using NAP5 or NAP10 (Pharmacia) using a PBSE buffer. The thiolated oligo (ES-1) is activated in TE buffer, pH 7.4 using 1 mM DDT and excess removed using spin column (BioRad). The activated streptavidin and oligonucleotide are preferably used directly in the cross linking reaction by mixing and incubation for about 1 hour in the dark at room temperature.

10 The modified streptavidin is preferably purified on a size-exclusion column or a anion-exchange column (MonoQ-HRS/5, Pharmacia).

Bifunctional complex molecules.

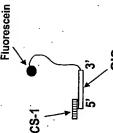
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Bifunctional complexes are preferably molecules that are composed of a nucleotide sequence that encodes for the displayed molecule. These complexes can be generated using various procedures, as disclosed elsewhere herein. The bifunctional complexes preferably contain an oligonucleotide that can be ligated or otherwise connected to the oligonucleotide sequence on the target mediated by the binding of the display molecule to the target.

This example describes the bifunctional complex as the IS-1 sequence which is labelled with a biotin in the 3'end (designated Y in the sequence). This oilgonucleotide is synthesized using Biotin-dT (Glen Research, # 10-1038-95), which will function as the display molecule and the oilgonucleotide sequence encoding the biotin moiety. The displayed blotin molecule has high affinity for streptavidin which will bring the coding oligonucleotide (CS-1) in close proximity of the identifier sequence (IS-1). This will promote the ligation between the ES-1 and the IS-1 oligonucleotides as shown below. The ligation is medated by the connector sequence (CS-1).

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also present. The flurorescein will not bind to the target molecule (strepta-A different bifunctional complex with a Fluorescein as display molecule is vidin) resulting in no proximity ligation between the ES-1 and the IS-1 oligonucleotides.

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Selection through proximity ligation

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ide. However, the ligation region is identical to the IS-1 oligonucleotide allowides longer permitting the distinction from the IS-1 oligonucleotide by running tures were incubated in 50 mM KCl, 10 mM Tris-HCl. pH 8.3, 1.5 mM MgCl₂, an agorose gel and determining the length of the oligonucleotides. The mixmolecule is encoded by another unique sequence in the CIS-1 oligonucleong ligation if proximity is achieved. The CIS-1 oligonucleotide is 10 nucleodisplay molecule (Fluorescein) included in the synthesis of the oligonucleomolecule to allow binding of the biotin molecule to streptavidin. Another bifunctional complex (CIS-1) was used as a control (100 pM) with a different 0.15 mM ATP, pH 7.4 for 1 hour to allow association of the Biotin (or Fluo-20 pM bifunctional complex (IS-1) is mixed with 100 pM conjugated target ide as a Fluorescein-dT (Glen Research, # 10-1056-95). This displayed rescein) to streptavidin.

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promote the connection between the ES-1 and IS-1 oligonucleotides together with 2 U T4 DNA ligase to start the ligation. The reaction was keep at 30°C The connector sequence (CS-1) is then added at 400 nM concentration to

33

WO 2005/026387

PCT/DK2004/000630

for 5 min and then 80°C for 20 min. The relatively high concentration of the connector oligonucleotide will saturate all IS-1 that have not been brought into proximity of an encoding sequence.

chain reaction (PCR) using primers corresponding to the 5'-end of the encodconsisted of an initial denaturation step of 94°C for 2 minutes followed by 20-ATTT). The primers are design to only amplify the ligated product. PCR was and 10 pmol of each primer in a reaction volume of 25 µl. The PCR reaction utes at 72°C was included. The PCR products were resolved by agarose gel performed using Ready-To-Go (RTG) PCR beads (Amersham Biosciences) ng sequence (PR-1: 5'-TAGTC GATGT AGCTA GCTAG) and the 3'-end of the identifier oligonucleotide sequence (PR-2: 5'-CAGCT TAGAC GGTAG minute and extension at 72°C for 1 minute. A final extension step of 2 minelectrophoresis and the band corresponding to the expected size was cut 45 cycles of denaturation at 94°C for 30 seconds, annealing at 58°C for 1 Amplification of the ligated products was performed with the polymerase from the gel and purified using QIAquick Gel Extraction Kit (QIAGEN). S 5 9

1014 different molecules. The same approach is used when screening library versus library. In this case the targets are encoded by different sequences plexes. The same approach can be used for larger libraries, at least up to The different length of the Biotin and Fluorescein identifier oligonucleotide sequences (IS-1 and CIS-1) is used to verify that the bifunctional complex This example describes a selection using two different bifunctional comwith Biotin have been enriched through the binding to streptavidin. but with identical coupling area.

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Cloning/sequencing

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TOP10 E. coli cells (Invitrogen) using standard procedures. The cells were cloned into the pCR4-TOPO vector (Invitrogen) according to the manufacturer's instructions. The resulting mixture was used for transformation of To sequence Individual PCR fragments the purified PCR products were

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124

beads and 5 pmol each of M13 forward and reverse primers according to the manufacturer's instructions. A sample of each PCR product was then treated plated on growth medium containing 100 µg/ml ampicillin and left at 37°C for wells containg 50 µl water. These wells were then boiled for 5 minutes and 20 µl mixture from each well was used in a PCR reaction using RTG PCR DYEnamic ET cycle sequencing kit (Amersham Biosciences) according to move degrade single stranded DNA and dNTPs and sequenced using the MegaBace 4000 capillary sequencer (Amersham Biosciences). Sequence 12-16 hours. Individual E.coli clones were picked and transferred to PCR with Exonuclease I (USB) and Shrimp Alkaline Phosphatase (USB) to rethe manufacturer's instructions and the reactions were analyzed on a outputs were analyzed with ContigExpress software (Informax Inc.).

Example 2: Library versus library

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genotype is associated with the phenotype, can be used together with bifunc-A library of bifunctional complexes is screened against another library of enscribes the use of bifunctional complexes together with mRNA displayed targets to perform library versus library screening. Any other library, where the coded peptides or proteins. Examples of other encoded peptides are ribosome displayed peptides or mRNA displayed peptides. This example deional complexes as described in this invention.

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A freshly transcribed mRNA (0.5 - 2.5 nmol) is prepared from an appropriate NaCl) by heating to 85° C for 30 sec followed by cooling to 4° C in 5 min. 100 library. The transcribed mRNA library is hybridized to biotinylated puromycinlinker (about 0.5 nmol) in 300 µl binding buffer (30 mM Tris, pH 7.0, 250 mM remove the liquid phase. The moist beads were then irradiated for 15 min at ul pre-washed Neutravidin beads (Pierce) is then added to the hybridization mixture and incubated for 30 min at 4 °C under rocking. Subsequently, the beads are washed in 3x 100 µl binding buffer followed by centrifugation to quently, the beads are washed with 100 µl plain water to yield the photoroom temperature with a 25W UV-lamp (Pyrex-filter, $\lambda > 300$ nm). Subse-

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WO 2005/026387

PCT/DK2004/000630

Res. 2000, 28:83). The puromycln-linker is also prepared according to Kurz protein fusions library according to the literature (Kurz et la., Nucleic Acids peptide fusion formation in rabbit reticulocyte lysate to produce the mRNAcrosslinked mRNA-puromycin template which is directly used for mRNA-

- amino group (50 µM) and the photo-cleavable biotin-reagent (NHS-PC-Biotin, et al. and biotinylated by carbamate bond formation between the puromycin 5 mM, EZ-Link TM-Biotin, Pierce Chemicals) in 25% DMSO/water for 2 h at room temperature followed by NaCI/EtOH precipitation.
- plementary to the distal region of the mRNA and which is able to promote an then used in the selection procedure to couple the target oligonucleotide with contain a coupling area. The coupling area is formed by a primer partly comextension on the mRNA strand with the coupling area. This coupling area is the identifier oligonucleotide mediated by the binding of displayed molecules The mRNA-peptide fusion library is then converted by a suitable primer to to the mRNA-peptide fusions. 9 5

1-100 pmol bifunctional complex library molecules are mixed with 1-100 pmol mRNA-peptide fusion library in a binding buffer (50 mM KCI, 10 mM Tris-HCI,

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the displayed molecules to the mRNA-peptide fusion molecules. The connecigase is added to start the ligation. The reaction is kept at 30°C for 5 min and then 80°C for 20 min. Amplification of the ligated products is performed using pH 8.3, 1.5 mM MgCl₂, 0.15 mM ATP. pH 7.4) for 1 hour to allow binding of tor sequence (an oligonucleotide the is complementary to the coupling area is then added at about 400 nM concentration to promote the connection between the target and identifier oligonucleotides. Subsequently, 2 U T4 DNA end of the target oligonucleotide and the 3'-end of the identifier oligonucleoa polymerase chain reaction (PCR) using primers corresponding to the 5'-

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formed using Ready-To-Go (RTG) PCR beads (Amersham Biosciences) and sisted of an initial denaturation step of 94°C for 2 minutes followed by 20-45 tide. The primers are design to only amplify the ligated product. PCR is per-10 pmol each primer in a reaction volume of 25 µl. The PCR reaction con-ဓ

cycles of denaturation at 94°C for 30 seconds, annealing at 58°C for 1 minute and extension at 72°C for 1 minute. A final extension step of 2 minutes at 72°C was Included. The PCR products are resolved by agarose gel electrophoresis and the band corresponding to the expected size is cut from the gel and purified using QIAquick Gel Extraction Kit (QIAGEN).

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The invention has been described in detail with reference to preferred embodiments thereof. However, it will be appreciated that those skilled in the art, upon consideration of this disclosure, may make modifications and improvements within the spirit and scope of the invention.

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WO 2005/026387

127

PCT/DK2004/000630

Claims

 A method for identifying display molecule(s) having affinity towards molecular target(s), comprising the steps of mixing one or more molecular target(s) associated with target oligonucleotide(s) and a library of bifunctional complexes, each bifunctional complex of the library comprising a display molecule attached to an identifier oligonucleotide, which codes for said display molecule,

coupling to the target oligonucleotide(s) the identifier oligonucleotide of complexes comprising display molecules binding to the target, and

deducing the identity of the binding display molecule(s) and/or the molecular target(s) from the coupled product between the identifier oligonucleotide(s) and the target oligonucleotide(s).

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 The method of claim 1, wherein the display molecule is a reaction product of two or more chemical entities and the identifier oligonucleotide comprises codons identifying the chemical entities.

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- The method of claim 1, wherein one or more members of the library are potentially binding compounds tagged with identifier oligonucleotides.
- 4. The method according to claim 1, 2 or, 3, wherein the chemical entities are precursors for a structural unit appearing in the display molecule.
- The method according to any of the claims 1 to 4, wherein some or all of the chemical entities are not naturally occurring α-amino acids or precursors thereof.

- The method according to claim 1 or 2, wherein each codon comprises 4 or more nucleotides.
- The method according to claim 1 or 2, wherein the display molecules of the library complexes are non-α-polypetides.
- The method according to claim 1 to 4, wherein the display molecules of the library complexes are non-nucleic acids.
- The method according to any of the preceding claims, wherein the
 display molecule has a molecular weight less than 2000 Dalton, preferably
 less than 1000 Dalton, and more preferred less than 500 Dalton.

128

- 10. The method according to any of the preceding claims, wherein the identifier oligonucleotide uniquely identifies the display molecule.
- more chemical entities are transferred to the nascent display molecule by a 11. The method according to any of the claims 1 to 10, wherein one or chemical building block further comprising an anti-codon.

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- anti-codon is transferred in conjunction with the chemical entity to the nas-12. The method according to claim 11, wherein the information of the sent complex.
- 13. The method according to any of the preceding claims, wherein the chemical entities are reacted without enzymatic interaction

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- 14. The method according to any of the claims 1 to 13, wherein the codons are separated by a framing sequence.
- 15. The method according to any of the claims 1 to 14, wherein the display molecule and the identifier oligonucleotide are joined by a selectively cleavable linker.

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- 16. The method according to claim 15, wherein the linker is cleaved by ir-
- 17. The method according to any of the claims, wherein the library comprises one, two or more different complexes
- 18. The method according to any of the claims 1 to 16, wherein the livary comprises 1,000 or more different complexes.

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19. The method according to claim 1, wherein the molecular target is of a

- 20. The method according to any of the claims 1 to 19, wherein the mobiological origin.
- 21. The method according to claim 20, wherein the target immobilized on he support forms a stable or quasi-stable dispersion. lecular target is immobilized on a solid support.

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- 22. The method according to claim 21, wherein a cleavable linker is present between the solid support and the molecular target.
- 23. The method according to any of the claims 1 to 22, wherein the moecular target is a protein 8

WO 2005/026387

PCT/DK2004/000630

24. The method according to claim 23, wherein the protein is selected rom the group consisting of kinases, proteases, phosphatases, and anti-

- 25. The method according to any of the clalms 1 to 24, wherein the mo
 - lecular target is a nucleic acid. S
- 26. The method according to claim 24, wherein the nucleic acid is an DNA or RNA aptamers.
- 27. The method according to any of the claims 23 to 26, wherein the target protein is attached to the nucleic acid responsible for the formation

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- ture step includes that a molecular target library comprising different peptides 28. The method according to any of the claims 1 to 27, wherein the mixeach attached to the nucleic acid responsible for the formation thereof is nixed with a library of complexes.
- 29. The method according to claim 28, wherein the library of complexes 30. The method according to any of the claims 1 to 29, wherein the target oligonucleotide is associated by a chemical synthesis to the molecular comprises a single bifunctional complex.
- 31. The method according to claim 30, wherein the molecular target is associated with the target oligonucleotide through one or more covalent or non-covalent bonds.

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- 32. The method according to any of the claims 1 to 31, wherein a bifunctional complex having a display molecule bInding to the molecular target
 - 33. The method according to clalm 32, wherein the display molecule is a constitutes the target oligonucleotIde associated with the molecular target. compound known to bind to the target.

- 34. The method according to claim 33, wherein a target is saturated with a known ligand prior to the mixing step.
- 35. The method according to claim 34, wherein the target oligonucleotide is associated with the molecular target during the mixing step. ဓ

plexes of a library of bifunctional complexes are associated with a common 36. The method according to claim 35, wherein two bifunctional commolecular target.

37. The method of claim 36, wherein the bifunctional complexes bind to the same binding site of the molecular target.

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- 38. The method of any of the claims 31 to 36, wherein the bifunctional complexes bind to discrete binding sites.
- 39. The method according to any of the preceding claims, wherein an inigeneration library, said second generation library being used in the method amended by reaction with one or more chemical entities to form a second tial ligand or a pool of ligands with potential affinity towards a target is

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according to any of the claims 1 to 39.

The method according to any of the preceding claims, wherein two or the method of claim 1, whereupon the identified display molecules binding to more targets interacting in a biological context separately are subjected to the two or more targets are linked via a suitable linker.,

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ing step and the target oligonucleotide identifies the molecular targets or the more molecular targets or type of molecular targets are involved in the mix-41. The method according to any of the claims 1 to 40, wherein two or type of molecular targets.

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- 42. The method according to any of the claims 1 to 41, wherein the mixing step includes the removal of non-binding library members prior to the coupling of the target oligonucleotide and the identifier oligonucleotide to-
- target oligonucleotide and/or the identifier oligonucleotide partly or fully is 43. The method according to any of the previous claims, wherein the hybridised to a complementing oligonucleotide.

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pling is performed using means selected from the group consisting of chemi-44. The method according to any of the claims 1 to 43, wherein the coucal means, enzymatic means, and design means.

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get oligonucleotide or a complementing target oligonucleotide and the identi-45. The method according to any of the claims 1 to 44, wherein the tar-

WO 2005/026387

PCT/DK2004/000630

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are joined together so as to allow for a polymerase to recognise the coupled fier oligonucleotide or a complementing identifier oligonucleotide operatively strand as a template.

- enzymatic means are selected from enzymes of the type polymerase, ligase 46. The method according to any of the preceding claims, wherein the and restriction enzyme, and any combination thereof.
- 47. The method according to claim 46, wherein a ligase is used to join the target oligonucleotide and the identifier oligonucleotide together.
- gion complementing a distal part of the identifier oligonucleotide Is used durng the coupling step so as to allow a ligase or a combination of a ligase and 48. The method of claim 47, wherein a connector oligonucleotide having a region complementing a distal part of the target oligonucleotide and a rea polymerase to join the identifier and target oligonucleotides together.

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- The method according to claim 48, wherein the ends of the oligonu
 - cleotides abut each other. 5
- 50. The method according to claim 49, wherein the region of the connecor oligonucleotide complementing a distal part of the identifier and/or target oligonucleotide is 6 to 16 nucleotides.
- 51. The method according to claims 49, wherein the region is 8 to 12 nu
 - cleotides. ನ
- 52. The method according to claim 48, wherein the connector oligonucleotide is added in excess.
- at the distal ends of the target and identifier oligonucleotides are complemen-53. The method according to any of the claims 1 to 46, wherein a region tary to each other and a polymerase is allowed to extend the target and/or the identifier oligonucleotide.

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- sticky end to allow a ligase or a polymerase or a mixture thereof to adjoin the 54. The method according to any of the claims 1 to 47, wherein the target oligonucleotide and/or the identifier oligonucleotide is provided with a oligonucleotides.
- 55. The method according to claim 54, wherein the sticky end Is formed by a restriction nuclease

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- get and the identifier oligonucleotide or sequences complementary thereto at 56. The method according to any of the claims 1 to 55, wherein the tarthe proximal end is provided with a priming site.
- get-display conjugate is recovered by chromatography following the coupling 57. The method according to any of the claims 1 to 56, wherein the tarof the target and the identifier oligonucleotides.

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- 58. The method according to claim 57, wherein the chromatography is size-exclusion chromatography.
- target oligonucleotide is amplified prior to decoding the identity of the display 59. The method according to claim 1, wherein the coupled identifier and

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- 60. The method according to any of the claim 1, wherein the coupled ollgonucleotide is amplified by PCR using priming sites positioned proximal to the display molecule and the molecular target, respectively.
- chemical moieties in each end of the coupled oligonucleotides are cleaved to 61. The method according to claim 59, wherein selective cleavable iberate the coupled oligonucleotide prior to amplification.

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- 62. The method according to any of the claims 1 to 61, wherein the coupled oligonucleotide is recovered and subjected to amplification.
- lached to an identifier oligonucleotide, which codes for said display molecule. gonucleotide and a bifunctional complex comprising a display molecule at-63. A conjugate comprising a molecular target associated with an oli-

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64. The conjugate of claim 63, wherein the display molecule is bound to the target.

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- gonucleotide and/or the identifier oligonucleotide are joined to the molecular arget and/or the display molecule, respectively, through a selectively cleav-65. The conjugate according to claims 63 or 64, wherein the target oli-
- 66. The conjugate according to any of the claims 63 to 65, wherein the target oligonucleotide is coupled to the identifier oligonucleotide.

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67. The conjugate according to claims 66, wherein the coupled oligonucleotide is amplifiable

WO 2005/026387

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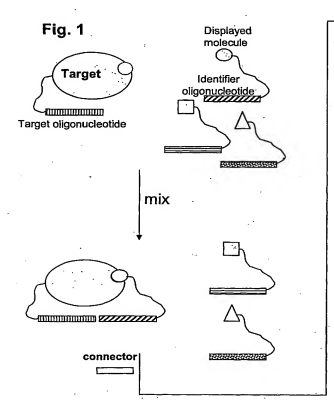
68. A display molecule identified by the method according to any of the claims 1 to 62

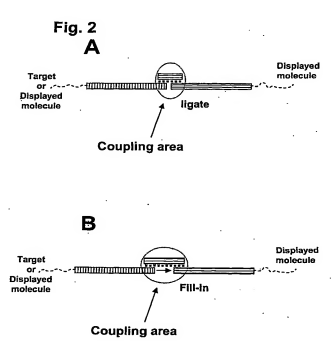
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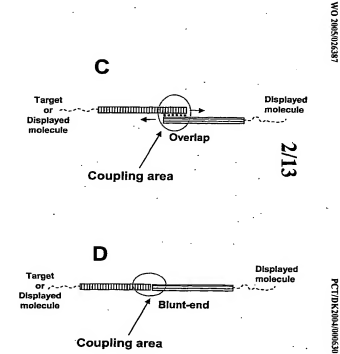
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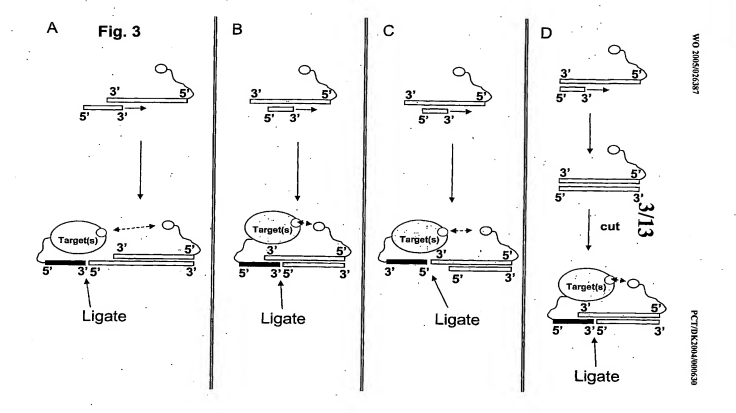
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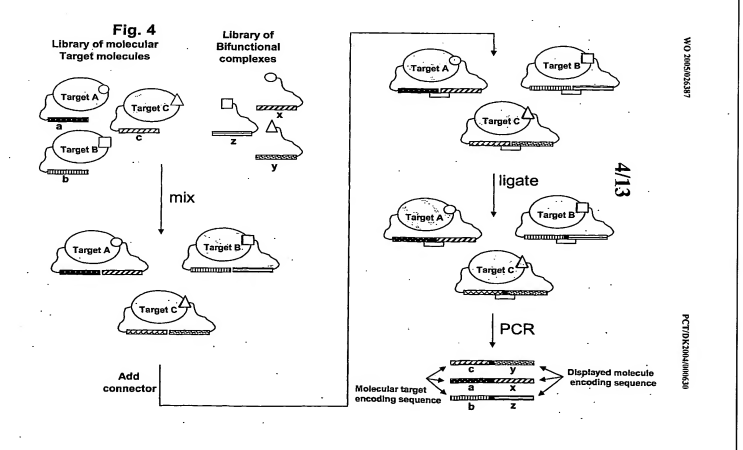
molecular target

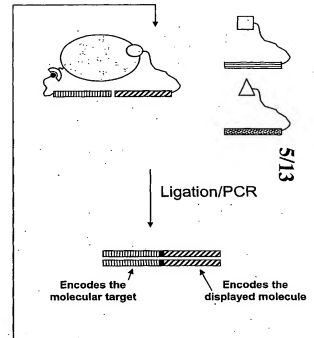


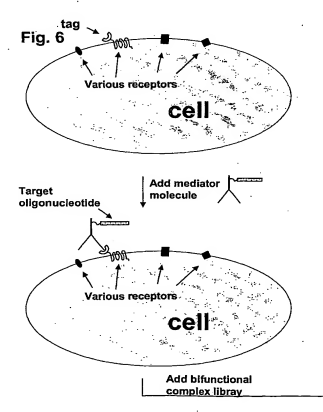


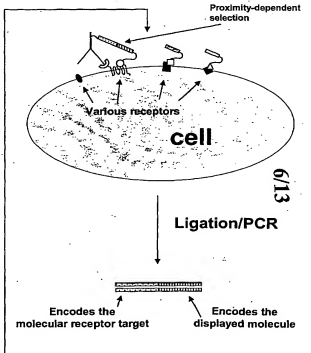


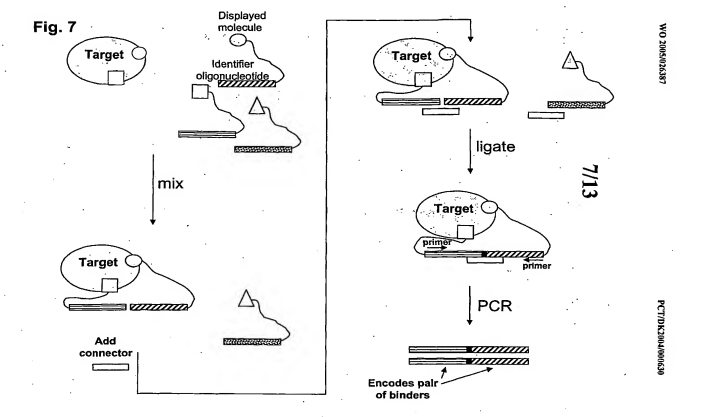


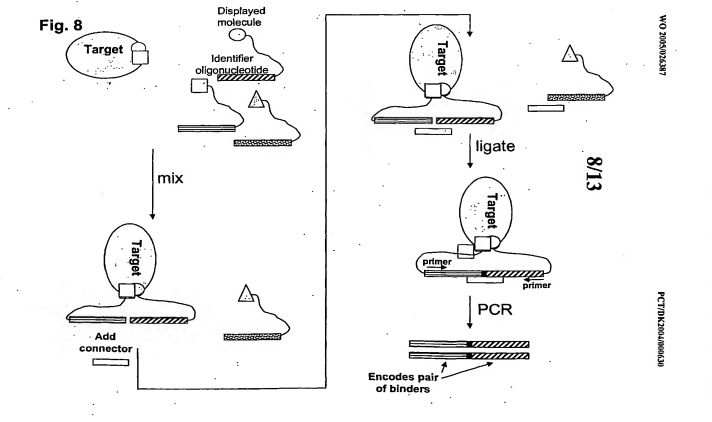


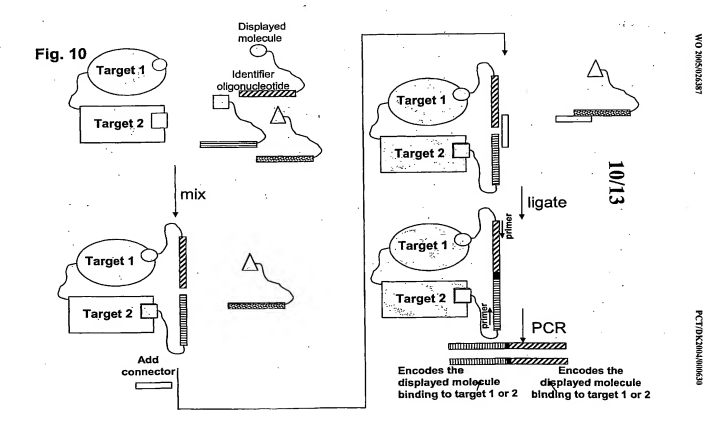












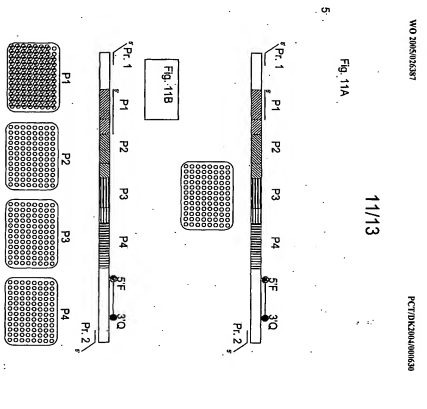
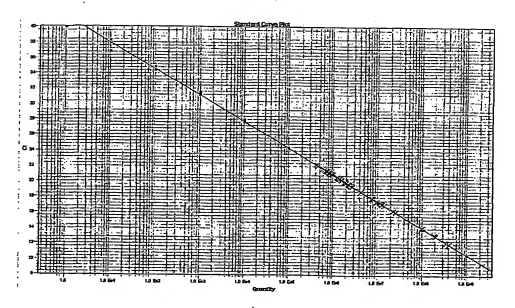


Fig. 12



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Fig. 13A

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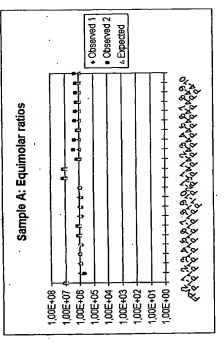
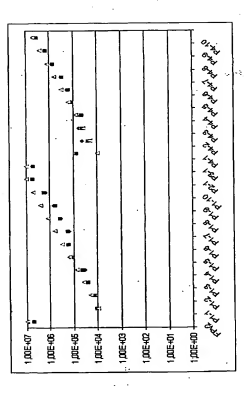


Fig. 13B Sample B



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